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[45] Date of Patent:

Dec. 22, 1998

[54] RAPID SUBCLONING USING SITE-SPECIFIC RECOMBINATION

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[73] Assignce: Baylor College of Medicine

[21] Appl. No.: 864,224

[22] Filed: Feb. 28, 1997

[56]

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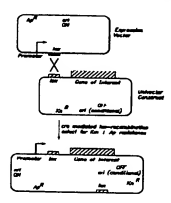
Primary Examiner—Johnny F. Railey, II Attorney, Agent, or Firm—Medlen & Carroll, 1.1.1

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ABSTRACT

The present invention provides compositions, including vectors, and methods for the rapid subcloning of nucleic acid sequences in vivo and in vitro. In particular, the invention provides vectors used to contain a gene of interest that comprise a sequence-specific recombinase target site. These vectors are used to rapidly transfer the gene of interest into any expression vector that contains a sequence-specific recombinase target site located downstream of a promoter element so that the gene of interest may be expressed.

24 Claims, 11 Drawing Sheets



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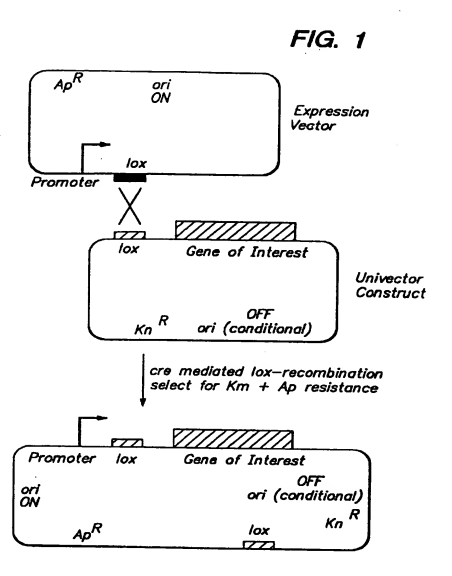


FIG. 2A HinD III (114) Bgl II (1896)NotI (401) PstI (1683) KpnI (409) EcoRI (452) XhoI (464) NdeI (472) PUNI 10 lox 2220 Бр Kan Nael (472) Ncol (479) BamHI (485) Notl (489) Hpal (503) Sall (515) Sacl (521) **BssHII** TERM (1335)Xbal (750) EcoRV (890)

FIG. 2B

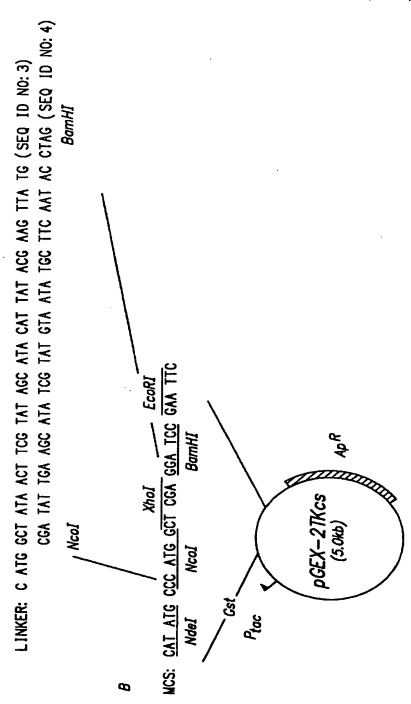
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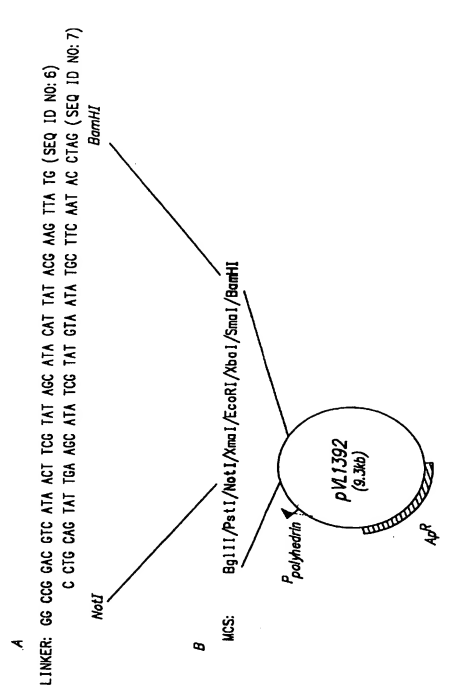
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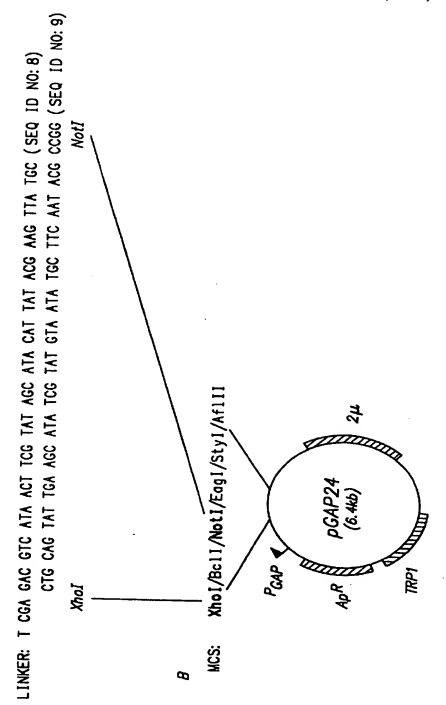
CONSTRUCTION OF pGst-lox:



CONSTRUCTION OF pW:1392-lox:

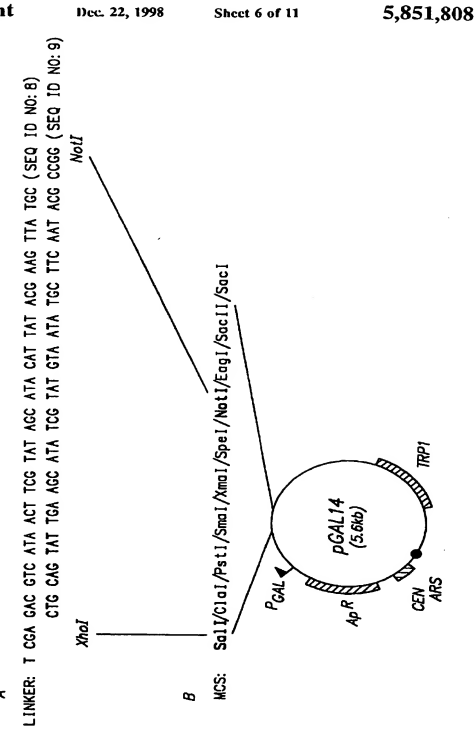


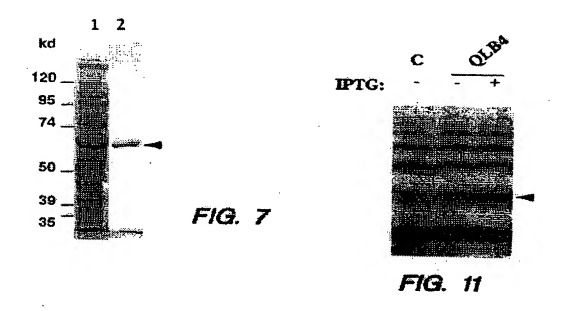
CONSTRUCTION OF pGAP24-lox:





CONSTRUCTION OF pGAL14-lox:





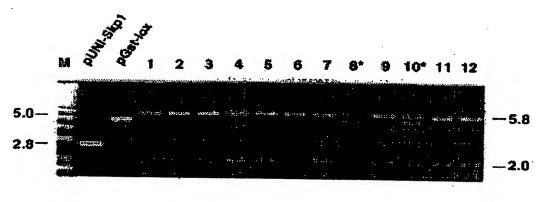
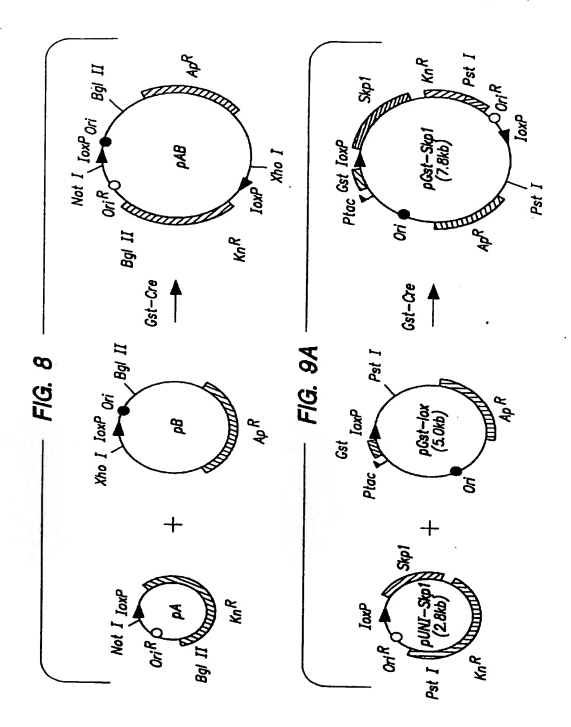
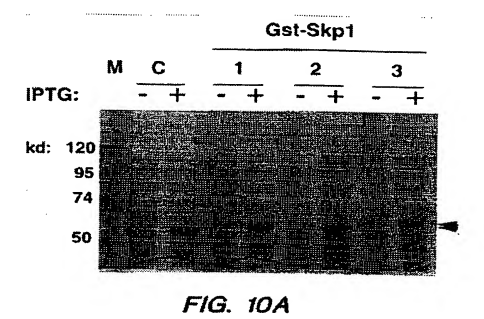


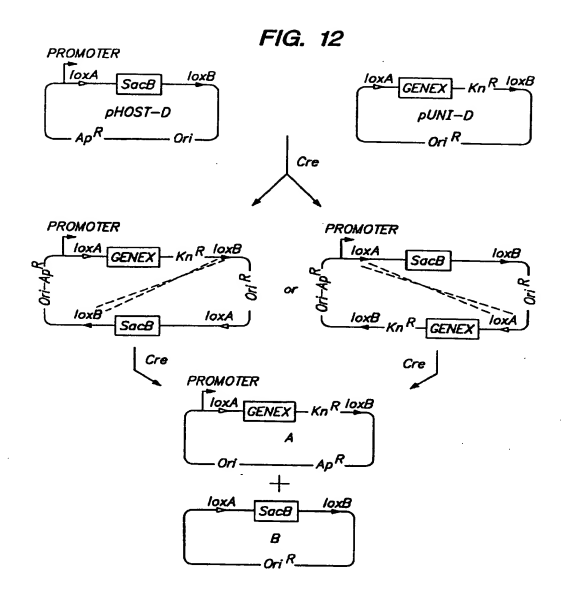
FIG. 9B





Gst-Skp1 IPTG:

FIG. 10B



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RAPID SUBCLONING USING SITE-SPECIFIC RECOMBINATION

FIELD OF THE INVENTION

The invention relates to recombinant DNA technology. In particular, the invention relates to compositions, including vectors, and methods for the rapid subcloming of nucleic acid sequences in vivo and in vitro.

BACKGROUND OF THE INVENTION

Molecular biotechnology has revolutionized the production of protein and polypeptide compounds of pharmacological importance. The advent of recombinant DNA technology permitted for the first time the production of proteins on a large scale in a recombinant host cell rather than by the laborious and expensive isolation of the protein from tissues which may contain minute quantities of the desired protein (e.g., isolation of human growth hormone from cadaver pituitary). The production of proteins, including human proteins, on a large scale in a beterologous host requires the ability to express the protein of interest in the heterologous host. This process typically involves isolation or cloming of the gene encoding the protein of interest followed by transfer of the coding region into an expression vector which contains elements (e.g., promoters) which direct the expression of the desired protein in the beterologous host cell. The most commonly used means of transferring or subcloning a coding region into an expression vector involves the in vitro use of restriction endomucleases and DNA ligases. Restriction endonucleases are enzymes which generally recognize and cleave a specific DNA sequence in a double-stranded DNA molecule. Restriction enzymes are used to excise the coding region from the cloming vector and the excised DNA fragment is then joined using DNA ligase to a suitably cleaved expression vector in such a manner that a functional protein may be expressed.

The ability to transfer the desired coding region to an expression vector is often limited by the availability or suitability of restriction enzyme recognition sites. Often an untiple restriction enzymes must be employed for the removal of the desired coding region and the reaction conditions used for each enzyme may differ such that it is necessary to perform the excision reactions in separate steps. In addition, it may be necessary to remove a particular enzyme used in an initial restriction enzyme reaction prior to completing all restriction enzyme digestions; this requires a time-consuming purification of the subcloning intermediate. Ideal methods for the subcloning of DNA molecules would permit the rapid transfer of the target DNA molecule from one vector to another in vitro or in vivo without the need to rely upon restriction enzyme digestions.

SUMMARY OF THE INVENTION

The present invention provides reagents and methods 35 which comprise a system for the rapid subcloning of nucleic acid sequences in vivo and in vitro without the need to use restriction enzymes. Accordingly, the present invention provides a nucleic acid construct comprising, in operable order:

a) a conditional origin of replication; b) a sequence-specific on recombinase target site having a 5' and a 3' end; and c) a unique restriction enzyme site, the restriction enzyme site located adjacent to the 3' end of the sequence-specific recombinase target site. The present invention is not limited by the nature of the conditional origin of replication 65 employed. A variety of conditional origins of replication, including temperature-sensitive replicons are known to the

art and may be employed on the nucleic acid constructs of the present invention. In a preferred embodiment, the nucleic acid construct further comprises a prokaryotic termination sequence. The present invention is not limited by the nature of the prokaryotic termination sequence chosen. In one embodiment, the prokaryotic termination sequence is the T7 termination sequence. A variety of termination sequences are known to the art and may be employed in the nucleic acid constructs of the present invention including, the T_{MY}, T_{LI}, T_{LI}

In another preferred embodiment, the nucleic acid construct further comprises a cukaryotic polyadenylation sequence. The present invention is not limited by the nature of the eukaryotic polyadenylation sequence chosen. In one embodiment, the eukaryotic polyadenylation sequence is selected from the group consisting of the bovine growth hormone polyadenylation sequence, the simian virus 40 polyadenylation sequence and the Herpes simplex virus thymidine kinase polyadenylation sequence. In yet another preferred embodiment, the nucleic soid construct further comprises a selectable marker gene. The present invention is not limited by the nature of the selectable marker gene chosen; the selectable marker may be a positive or negative selectable marker. In a preferred embodiment, the selectable marker is selected from the group consisting of the kanamycin resistance gene, the ampicillin resistance gene, the tetracycline resistance gene, the chloramphenicol resistance gene, the streptomycin resistance gene, the strA gene and the sacB gene.

The present invention is not limited by the nature of the sequence-specific recombinase target site employed on the nucleic acid construct. In one embodiment, the sequence-specific recombinase target site is selected from the group consisting of loxP, loxP2, loxP31, loxP31, loxB, loxC2, loxL, loxR, loxA86, loxA117,frt, dif, and att.

In a preferred embodiment, the nucleic construct further comprises a gene of interest inserted into the unique restriction enzyme site.

In a preferred embodiment, the nucleic acid construct has the nucleotide sequence set forth in SEQ ID NO:1 (i.e., pUNI-10).

The present invention further provides a nucleic acid construct comprising, in order 5' to 3': a) a conditional origin of replication; b) a sequence-specific recombinase target site; c) a polylinker; and d) a selectable marker gene. The present invention is not limited by the nature of the polylinker employed. Any collection of two or more restriction enzyme sites which are unique to the nucleic seid construct may be employed as the polylinker. The present invention is not limited by the nature of the conditional origin of replication employed. A variety of conditional origins of replication, including temperature-sensitive replicons are known to the art and may be employed on the aucleic acid constructs of the present invention. In a preferred embodiment, the nucleic acid construct further comprises a prokaryotic termination sequence. The present invention is not limited by the nature of the prokaryotic termination sequence chosen. In one embodiment, the prokaryotic termination sequence is the '17 termination sequence. In another preferred embodiment, the nucleic acid

construct further comprises a cukaryotic polyadenylation sequence. The present invention is not limited by the nature of the eukaryotic polyademylation sequence chosen. In one embodiment, the eukaryotic polyadenylation sequence is selected from the group consisting of the bovine growth hormone polyadenylation sequence, the simian virus 40 polyadenylation sequence and the Herpes simplex virus thymidine kinase polyadenylation sequence. In yet another preferred embodiment, the nucleic acid construct further comprises a selectable marker gene. The present invention is 10 not limited by the nature of the selectable marker gene chosen; the selectable marker may be a positive or negative selectable marker. In a preferred embodiment, the selectable marker is selected from the group consisting of the kanamycin resistance gene, the ampicillin resistance gene, the 15 tetracycline resistance gene, the chloramphenical resistance gene, the streptomycin resistance gene, the strA gene and the

The present invention is not limited by the nature of the sequence-specific recombinase target site employed on the 20 nucleic acid construct. In one embudiment, the sequence-specific recombinase target site is selected from the group consisting of loxP, loxP2, loxP3, loxP23, loxP511, loxB, loxC2, loxL, loxR, loxA86, loxA117,frt, dif, and att.

In a preferred embodiment, the nucleic construct further ²⁵ comprises a gene of interest inserted into the polylinker.

The present invention also provides a nucleic acid construct comprising in operable order: a) an origin of replication; b) a promoter element having a 5' and a 3 end; and c) a sequence-specific recombinase target site having a 5' and a 3' end. In a preferred embodiment the 3' end of the promoter element is located upstream of the 5' end of said a sequence-specific recombinase target site. The present invention is not limited by the nature of the origin of 35 replication employed. A variety of non-conditional origins of replication are known to the art and may be employed on the nucleic acid constructs of the present invention. The invention is not limited by the nature of the promoter element employed. Those skilled in the art know that the choice of the promoter element depends upon the type of host cell to be employed for expressing a gene(s) under the transcriptional control of the chosen promoter element. A wide variety of promoter elements functional in prokaryotic (c.g., E. coli) and cukaryotic (c.g., yeast, insect, marimals including humans) cells are known to the art and may be employed in the nucleic acid constructs of the present invention.

In another preferred embodiment, the nucleic acid construct further comprises a selectable marker gene. The 50 present invention is not limited by the nature of the selectable marker gene chosen; the selectable marker may be a positive or negative selectable marker. In a preferred embodiment, the selectable marker is selected from the group consisting of the kanamycin resistance gene, the 55 ampicillin resistance gene, the tetracycline resistance gene, the chloramphenical resistance gene, the streptomycin resistance gene, the strA gene and the sacB gene. The present invention is not limited by the nature of the sequencespecific recombinase target site employed on the nucleic acid construct. In one embodiment, the sequence-specific recombinase target site is selected from the group consisting of loxP, loxP2, loxP3, loxP23, loxP511, loxB, loxC2, loxL, loxR, loxA86, loxA117, frt, dif and att.

The present invention further provides a nucleic acid 65 construct comprising in operable order: a) a promoter element having a 5' and a 3 end; b) a first sequence-specific

recombinase target site having a 5' and a 3' end, wherein the 3' end of the promoter element is located upstream of the 5' end of the a sequence-specific recombinase target site; c) a gene of interest joined to the 3' end of the sequence-specific recombinase target site such that a functional translational reading frame is created; d) a conditional origin of replication; e) a first selectable marker gene; f) a second sequencespecific recombinase target site; and g) an origin of replication. In a preferred embodiment, the first and the second sequence-specific recombinase target site have the same nucleotide sequence (e.g., both comprise a loxP site). The first and the second sequence-specific recombinase target site need not have the same nucleotide sequence provided the different sites can recombine with one another. The present invention is not limited by the nature of either the conditional origin of replication or the non-conditional origin of replication employed.

In a preferred embodiment, the nucleic acid construct further comprises a second selectable marker gene. Preferably the first and the second selectable marker genes are different selectable marker genes. The present invention is not limited by the nature of the selectable marker genes chosen; the selectable marker genes may be positive or negative selectable marker genes. As discussed above, the present invention is not limited by the nature of the promoter element nor the nature of the sequence-specific recombinase target sites chosen.

The present invention also provides a method for the in vitro recombination of nucleic acid constructs, comprising: a) providing: i) a first nucleic acid construct comprising, in operable order, a conditional origin of replication, a first sequence-specific recombinase target site and a first selectable marker gene; ii) a second nucleic acid construct comprising, in operable order, an origin of replication, a promoter element and a second sequence-specific recombinase target site; and iii) a site-specific recombinase; b) contacting the first and the second constructs in vitro with the site-specific recombinase under conditions such that the first and second constructs are recombined to form a third nucleic acid construct.

In a preferred embodiment, the first construct employed in the method further comprises a gene of interest and the recombination of the first and second constructs places the gene of interest under the transcriptional control of the promoter element. In another preferred embodiment, the second construct further comprises a nucleotide sequence encuding an affinity domain and the recombination of the first and second constructs results in placing the gene of interest in frame with the sequence encuding the affinity domain. The present invention is not limited by the nature of the affinity domain employed; a variety of suitable affinity domains are known to the art including glutathione-Stransferase, the maltose binding protein, protein A, polyhistidine tracts, etc.

In a preferred embodiment, the second construct further comprises a second selectable marker gene, the second selectable marker being different from the first selectable marker (present on the first construct). The present invention is not limited by the nature of the selectable marker genes chosen; the selectable marker genes may be positive or negative selectable marker genes. As discussed above, the present invention is not limited by the nature of the promoter element nor the nature of the sequence-specific recombinase target sites chosen.

The present invention further provides a method for the recombination of nucleic acid constructs in a prokaryotic

host, comprising a) providing: i) a first nucleic acid construct comprising a conditional origin of replication, a first sequence-specific recombinase target site having a 5' and a 3' end, a unique restriction enzyme site, the restriction enzyme site located adjacent to the 3' end of the sequence- 5 specific recombinase target site, a first selectable marker gene and a eukaryotic polyadenylation sequence; ii) a second nucleic acid construct comprising an origin of replication, a promoter element having a 5' and a 3' end and a second sequence-specific recombinase target site baving a 10 5' and a 3' end; iii) a procaryotic bost cell expressing a site-specific recombinase; b) introducing the first and said second DNA constructs into the prokaryotic host cell under conditions such that the first and second DNA constructs are recombined to form a third nucleic acid construct capable of 15 replicating in the host cell.

In a preferred embodiment, the 3' end of the promoter element is located upstream of the 5' end of the second sequence-specific recombinase target site.

In another preferred embodiment, the method further 20 comprises growing the bost cell containing the third construct under conditions which select for the presence of the third construct.

In a preferred embodiment, the first construct further comprises a prokaryotic termination sequence. The present invention is not limited by the nature of the prokaryotic termination sequence chosen. In one embodiment, the prokaryotic termination sequence is the T7 termination sequence.

in another preferred embodiment, the first construct further comprises a gene of interest inserted into the unique restriction endonuclease recognition site.

The present invention also provides a fusion protein comprising protein sequences derived from a glutathione-S-transferase (Gst) protein and a Cre protein; the Gst-Cre fusion protein may be provided in a purified form. In a preferred embediment, the Gst-Cre fusion protein has the amino acid sequence of SEQ ID NO:11. In another preferred embediment, the Gst-Cre fusion protein is encoded by the nucleic acid sequence of SEQ ID NO:10.

DESCRIPTION OF THE DRAWINGS

FIG. 1 provides a schematic illustrating certain elements of the pUNI vectors and the Univector Fusion System.

FIG. 2A provides a schematic map of the pUNI-10 vector; the locations of selected restriction enzyme sites are indicated and unique sites are indicated by the use of bold type.

FIG. 2B shows the DNA sequence of the loxP site and the polylinkers contained within pUNI-10 (i.e., nucleotides 401-530 of SEQ ID NO:1).

FIG. 3A shows the oligonucleotides (SEQ ID NOS:4 and 5) which were annealed to insert a loxP site into the polylinker of pGEX-2TKes to create pgst-lox.

FIG. 3B provides a schematic map of pGEX-21Kes 55 which includes an enlargement of the multiple cloning site (MCS).

FIG. 4A shows the oligonucleotides (SEQ ID NOS:6 and 7) which were annealed to insert a loxP site into the polylinker of pVL1392 to create pVL1392-lox.

FIG. 4H provides a schematic map of pVI.1392 which includes an enlargement of the multiple choing site (MCS); the ampicillin resistance gene (Ap®) and the tac promoter (P_{ma}) are indicated.

FIG. 5A shows the oligonucleotides (SEO ID NOS:8 and 65 9) which were annealed to insert a loxP site into the polylinker of pGAP24 to create pGAP24-lox.

FIG. 5H provides a schematic map of pGAP24 which includes an enlargement of the multiple cloning site (MCS); the ampicillin resistance gene (Ap®), the GAP promoter (P_{tRP}), the origin from the 2 μ m circle (2 μ) and the TRPI gene, encoding N-(5'-phosphoribosyl)-anthrapilate synthetase, (TRPI) are indicated.

FIG. 6A shows the oligonucleotides (SEQ ID NOS:8 and 9) which were annealed to insert a loxP site into the polylinker of pGAL14 to create pGAL14-lox.

FIG. 6B provides a schematic map of pGAI.14 which includes an enlargement of the multiple cloning site (MCS); the ampicillin resistance gene (Ap \oplus), the GAL promoter (P_{GAI.}), the yeast centromeric sequences (CEN), yeast autonomous replication sequences (ARS) and the TRPI gene (IRII) are indicated.

FIG. 7 shows a Coomassic blue-stained SDS-PAGE gel showing the purification of Gst-Cre from E. coli cells containing pQL123.

FIG. 8 provides a schematic showing the strategy employed for the in vitro recombination of a pUNI vector ("pA," pUNI-5) with a pIIOST vector ("pB," pQL103) to create a fuscal construct ("pAB"). The relevant markers on each construct are indicated as are selected restriction enzyme sites.

FIG. 9A provides a schematic showing the starting constructs (pUNI-Skpl and pgst-lox) and the predicted fusion construct (p(kt-Skpl) generated by an in vitro fusion reaction.

FIG. 9B shows an ethicium bromide-stained gel is showing the separation of restriction fragments generated by the digestion of pUNI-Skpl, pgst-lox and pCst-Skpl.

FIG. 10A shows a Coomassic blue-stained SDS-PAGE gel showing the expression of the Gst-Skpl protein from E. culi cells containing pGst-Skpl.

FIG. 10B shows a Western blot of an SDS-PAGE get containing extracts prepared from E. coli cells containing pGst-Skpl which was probed using an anti-Skpl antibody.

FIG. 11 shows a Western blot of an SIS-PAGE gel containing extracts prepared from E. coll cells (QLR4) containing either a conventionally constructed Gst-Skpt plasmid or pCist-Skpl (produced by an in vitro fusion reaction).

FIG. 12 provides a schematic illustrating the in vivo gene trap method for the recombination of lox-containing vectors in a host cell constitutively expressing the Cre protein.

FIG. 13 provides the nucleotide sequence of the wild-type loxP site (SEQ ID NO:12), the loxP2 site (SEQ ID NO:13), the loxP3 site (SEQ ID NO:14) and the loxP23 site (SEQ ID NO:15).

DEFINITIONS

To facilitate understanding of the invention, a number of terms are defined below.

As used herein, "a conditional origin of replication" refers to an origin of replication that requires the presence of a functional trans-acting factor (e.g., a replication factor) in a prokaryotic host cell. Conditional origins of replication encompass temperature-sensitive replicons such as reppSC101".

As used berein, the term "origin of replication" refers to an origin of replication that is functional in a broad range of prokaryotic bost cells (i.e., a normal or non-conditional origin of replication such as the ColEl origin and its derivatives). The terms "sequence-specific recombinase" and "sitespecific recombinase" refer to enzymes that recognize and hind to a short nucleic acid site or sequence and catalyze the recombination of nucleic acid in relation to these sites.

The terms "sequence-specific recombinase target site" 5 and "site-specific recombinase target site" refer to short nucleic acid site or sequence which is recognized by a sequence- or site-specific recombinase and which become the crossover regions during the site-specific recombinase target to sites include, but are not limited to, lox sites, frt sites, att sites and dif sites.

The term "lox site" as used herein refers to a nucleotide sequence at which the product of the cre gene of hacteriophage Pl, Cre recombinase, can catalyze a site-specific recombination. A variety of lox sites are known to the art including the naturally occurring loxP (the sequence found in the Pl genome), loxB, loxL and loxR (these are found in the E. coli chromosome) as well as a number of mutant or variant lox sites such as loxP511, loxA86, loxA117, loxC2, loxP2, loxP3 and loxP23.

The term "frt site" as used herein refers to a nucleotide sequence at which the product of the FLP gene of the yeast 2 μ m plasmid, FLP recombinase, can catalyze a site-specific recombination.

The term "unique restriction enzyme site" indicates that the recognition sequence for a given restriction enzyme appears once within a nucleic acid molecule. For example, the EcoRI site is a unique restriction enzyme site within the plasmid pUNI-10 (SEQ ID NO:1).

A restriction enzyme site is said to be located "adjacent to the 3' end of a sequence-specific recombinase target site" if the restriction enzyme recognition site is located downstream of the 3' end of the sequence-specific recombinase target site. The adjacent restriction enzyme site may, but need not, be contiguous with the last or 3' nucleotide comprising the sequence-specific recombinase target site. For example, the EcoRI site of pUNI-10 is located adjacent (within 3 nucleotides) to the 3' end of the loxP site (see PIG. 2B); the Xhol, Ndel, Neel sites are also adjacent (ie., within about 10-150 nucleotides) to the loxP site but these sites are not contiguous with the 3' end of the loxP site in pUNI-10.

The terms "polylinker" or "multiple cloning site" refer to a cluster of restriction enzyme sites on a nucleic acid 45 construct which are utilized for the insertion and/or excision of nucleic acid sequences such as the coding region of a gene, lox sites, etc.

The term "prokaryotic termination sequence" refers to a nucleic acid sequence which is recognized by the RNA so nucleic acid sequence which is recognized by the RNA so polymerase of a prokaryotic host cell and results in the termination of transcription. Prokaryotic termination sequences commonly comprise a GC-rich region that has a twofold symmetry followed by an AT-rich sequence [Stryer, supra]. A commonly used prokaryotic termination sequence is the T7 termination sequence. A variety of termination sequences are known to the art and may be employed in the nucleic acid constructs of the present invention including, the T_{MT}, T_{L1}, T_{L2}, T_{L2}, T_{R1}, T_{R2}, T_{AC}-termination signals derived from the bacteriophage lambda [Lambda II, Hendrix et al. Eds., supra] and termination signals derived from bacterial genes such as the trp gene of E. coli [Stryer, supra].

The term "eukaryotic polyadenylation sequence" (also referred to as a "poly A site" or "poly A sequence") as used herein denotes a DNA sequence which directs both the 65 termination and polyadenylation of the nascent RNA transcript. Efficient polyadenylation of the recombinant transcript.

script is desirable as transcripts lacking a poly \wedge tail are unstable and are rapidly degraded. The poly \wedge signal utilized in an expression vector may be "heterologous" or "endogcoous." An endogenous poly A signal is one that is found naturally at the 3 end of the coding region of a given gene in the genome. A heterologous poly A signal is one which is one which is isolated from one gene and placed 3' of another gene. A commonly used heterologous poly A signal is the SV40 poly A signal. The SV40 poly A signal is contained on a 237 bp BamHI/BcII restriction fragment and directs both termination and polyadenylation [J. Sambrook, supra, at 16.6-16.7]; numerous vectors contain the SV40 poly A signal [e.g., pCEP4, pREP4, pEBVHis (Invitrogen)]. Another commonly used heterologous poly A signal is derived from the bovine growth hormone (BGII) gene; the BGII poly A signal is available on a number of commercially available vectors [e.g., pcDNA3.1, pZcoSV2, pSccTag (Invitrogen)]. The poly A signal from the Herpes simplex virus thymidine kinase (IISV tk) gene is also used as a poly A signal on expression vectors; vectors containing the HSV tk poly A signal include the pBK-CMV, pBK-RSV, pOP13CAT vectors from Stratagene.

As used herein, the terms "selectable marker" or "selectable marker gene" refers to the use of a gene which encodes an enzymatic activity that confers the ability to grow in medium lacking what would otherwise be an essential nutrient (e.g., the TRPI gene in yeast cells); in addition, a selectable marker may confer resistance to an antibiotic or drug upon the cell in which the selectable marker is expressed. A selectable marker may be used to confer a particular phonotype upon a host cell. When a bost cell must express a selectable marker to grow in selective medium, the marker is said to be a positive selectable marker (e.g., antibiotic resistance genes which confer the ability to grow in the presence of the appropriate antibiotic). Selectable markers can also be used to select against bost cells containing a particular gene (e.g., the sacB gene which, if expressed, kills the bacterial host cells grown in medium containing 5% sucrose); selectable markers used in this manner are referred to as negative selectable markers or counter-selectable markers.

As used herein, the term "vector" is used in reference to nucleic arid molecules that transfer DNA segment(s) from one cell to another. The term "vehicle" is sometimes used interchangeably with "vector." A "vector" is a type of "nucleic acid construct." The term "nucleic acid construct includes circular nucleic acid constructs such as plasmid constructs, plasmid constructs, cosmid vectors, etc. as well as linear nucleic acid constructs (e.g., \(\lambda\) phage constructs, PCR products). The nucleic acid construct may comprise expression signals such as a promoter and/or an enhancer (in such a case it is referred to as an expression vector).

The term "expression vector" as used herein refers to a recombinant DNA molecule containing a desired coding sequence and appropriate nucleic acid sequences necessary for the expression of the operably linked coding sequence in a particular host organism. Nucleic acid sequences necessary for expression in procuryous usually include a promoter, an operator (optional), and a ribosome binding site, often along with other sequences. Eucaryotic cells are known to utilize promoters, enhancers, and termination and polyadenylation signals.

The terms "in operable combination", "in operable order" and "operably linked" as used herein refer to the linkage of nucleic acid sequences in such a manner that a nucleic acid molecule capable of directing the transcription of a given gene and/or the synthesis of a desired protein molecule is

produced. The term also refers to the linkage of amino acid sequences in such a manner so that a functional protein is produced.

The terms "transformation" and "transfection" as used herein refer to the introduction of foreign DNA into prokaryotic or eucaryotic cells. Transformation of prokaryotic cells may be accomplished by a variety of means known to the art including the treatment of host cells with CaCl₂ to make competent cells, electroporation, etc. Transfection of eukaryotic cells may be accomplished by a variety of means known to the art including calcium phosphate-DNA co-precipitation, DEAE-dextran-mediated transfection, polybrene-mediated transfection, electroporation, microinjection, liposome fusion, lipofection, protoplast fusion, retroviral infection, and biolistics.

As used herein, the terms "restriction endonucleases" and "restriction enzymes" refer to bacterial enzymes, each of which cut double-stranded DNA at or near a specific nucleotide sequence.

As used herein, the term "recombinant DNA molecule" as 20 used herein refers to a DNA molecule which is comprised of segments of DNA joined together by means of molecular biological techniques.

DNA molecules are said to have "5' ends" and "3' ends" because mononucleotides are reacted to make oligonucle- 25 otides in a manner such that the 5' phosphate of one mononucleotide pentose ring is attached to the 3' oxygen of its neighbor in one direction via a phosphodiester linkage. Therefore, an end of an oligonucleotides referred to as the "5' end" if its 5' phosphate is not linked to the 3' oxygen of a mononucleotide pentose ring and as the "3' end" if its 3' oxygen is not linked to a 5' phosphate of a subsequent mononucleotide pentose ring. As used herein, a nucleic acid sequence, even if internal to a larger oligonucleotide, also may be said to have 5' and 3' ends. In either a linear or 35 circular DNA molecule, discrete elements are referred to as being "upstream" or 5' of the "downstream" or 3' elements. This terminology reflects the fact that transcription proceeds in a 5' to 3' fashion along the DNA strand. The promoter and enhancer elements which direct transcription of a linked 40 gene are generally located 5' or upstream of the coding region. However, enhancer elements can exert their effect even when located 3' of the promoter element and the coding region. Transcription termination and polyadenylation sig-nals are located 3' or downstream of the coding region.

The 3' end of a promoter element is said to be located upstream of the 5' end of a sequence-specific recombinase target site when (moving in a 5' to 3' direction along the nucleic acid molecule) the 3' terminus of a promoter element (the transcription start site is taken as the 3' end of a 50 promoter element) precedes the 5' end of the sequencespecific recombinase target site. The 3' end of the promoter element may be located adjacent (generally within about 0 to 500 hp) to the 5' end of the sequence-specific recombinate target site; such an arrangement is used when the pHOST 55 ctor is not intended to permit the expression of a translational fusion with the gene of interest donated by a pUNI voctor. Alternatively when the pHOST vector is intended to permit the expression of a translational fusion, the 3' end of the promoter element is located upstream of both the so sequences encoding the amino-terminus of a fusion protein and the 5' end of the sequence-specific recombinase target site; in this case, the 5' end of the sequence-specific recombinase target site is located within the coding region of the fusion protein (e.g., located downstream of both the pro-moter element and the sequences encoding the affinity domain, such as Gst).

As used herein, the term "an oligonucleotide having a nucleotide sequence encoding a gene" means a nucleic acid sequence comprising the coding region of a gene or in other words the nucleic acid sequence which encodes a gene product. The coding region may be present in either a cDNA, genomic DNA or RNA form. When present in a DNA form, the oligonucleotide may be single-stranded (i.e., the sense strand) or double-stranded. Suitable control elements such as enhancers/promoters, splice junctions, polyadenylation signals, etc. may be placed in close proximity to the coding region of the gene if needed to permit proper initiation of transcription and/or correct processing of the primary RNA transcript. Alternatively, the coding region utilized in the vectors of the present invention may contain endogenous enhancers/promoters, splice junctions, intervening sequences, polyadenylation signals, etc. or a combination of both endogenous and exogenous control elements.

As used herein, the term "regulatory element" refers to a genetic element which controls some aspect of the expression of nucleic acid sequences. For example, a promoter is a regulatory element which facilitates the initiation of transcription of an operably linked coding region. Other regulatory elements are splicing signals, polyadenylation signals, termination signals, etc. (defined infra).

Transcriptional control signals in cultaryotes comprise "promoter" and "enhancer" elements. Promoters and enhancers consist of short arrays of DNA sequences that interact specifically with cellular proteins involved in transcription [Maniatis, T. et al., Science 236:1237 (1987)]. Promoter and enhancer elements have been isolated from a variety of cuearyotic sources including genes in yeast, insect and mammalian colls and viruses (analogous control elements, i.e., promoters, are also found in procaryotes). The selection of a particular promoter and enhancer depends on what cell type is to be used to express the protein of interest. Some eucaryotic promoters and enhancers have a broad host range while others are functional in a limited subset of cell types [for review see Voss, S.D. et al., Trends Biochem. Sci. 11:287 (1986) and Maniatis, T. et al., supra (1987)]. For example, the SV40 early gene enhancer is very active in a wide variety of cell types from many mammalian species and has been widely used for the expression of proteins in mammalian cells [Dijkema, R. et al., EMBO J. 4:761 (1985)]. Two other examples of promoter/enhancer elements active in a broad range of mammalian cell types are those from the human elongation factor 10 gene [Uetsuki, T. et al., J. Blok Chem., 264:5791 (1989), Kim, D. W. et al., Gene 91:217 (1990) and Mizushima, S. and Nagata, S., Nuc. Acids. Res., 18:5322 (1990)] and the long terminal repeats of the Rous sarcoma virus [Gorman, C. M. et al., Proc. Natl. Acad. Sci. USA 79:6777 (1982)] and the human cytomegalovirus [Boshart, M. et al., Cell 41:521 (1985)].

As used berein, the term "promoter/enhancer" denotes a segment of INA which contains sequences capable of providing both promoter and enhancer functions (i.e., the functions provided by a promoter element and an enhancer clement, see above for a discussion of these functions). For example, the long terminal repeats of retroviruses contain both promoter and enhancer functions. The enhancer/promoter may be "endogenous" or "exogenous" or "heterologous." An "endogenous" enhancer/promoter is one which is naturally linked with a given gene in the genome. An "exogenous" or "heterologous" enhancer/promoter is one which is placed in juxtaposition to a gene by means of genetic manipulation (i.e., molecular biological techniques) such that transcription of that gene is directed by the linked enhancer/promoter.

The presence of "splicing signals" on an expression vector often results in higher levels of expression of the recombinant transcript. Splicing signals mediate the removal of introns from the primary RNA transcript and consist of a splice donor and acceptor site [Sambrook, J. et al., Molecular Cloning: A Laboratory Manual, 2nd ed., Cold Spring Harbor Laboratory Press, New York (1989) pp. 16.7–16.8]. A commonly used splice donor and acceptor site is the splice junction from the 16S RNA of SV40.

Efficient expression of recombinant DNA sequences in 10 eucaryotic cells requires expression of signals directing the efficient termination and polyadenylation of the resulting transcript. Transcription termination signals are generally found downstream of the polyadenylation signal and are a few hundred nucleotides in length. The term "poly A site" or "poly A sequence" as used herein denotes a DNA sequence which directs both the termination and polyadenylation of the nascent RNA transcript. Efficient polyadenylation of the recombinant transcript is desirable as transcripts lacking a poly A tail are unstable and are rapidly degraded. The poly 20 A signal utilized in an expression vector may be "heterologous" or "endogenous." An endogenous poly A signal is one that is found naturally at the 3' end of the coding region of a given gene in the genome. A beterologous poly A signal is one which is one which is isolated from one gene and placed 25 3' of another gene.

Eucaryotic expression vectors may also contain "viral replicons" or "viral origins of replication." Viral replicons are viral DNA sequences which allow for the extrachromosomal replication of a vector in a bost cell expressing the appropriate replication factors. Vectors which contain either the SV40 or polyoma virus origin of replication replicate to high copy number (up to 10° copies/cell) in cells that express the appropriate viral T antigen. Vectors which contain the replicons from bovine papillomavirus or Epstein-Barr virus replicate extrachromosomally at low copy number (-100 copies/cell).

As used herein, the terms "nucleic acid molecule encoding," "DNA sequence encoding," and "DNA encoding" refer to the order or sequence of deoxyribonucleotides along a strand of deoxyribonucleic acid. The order of these deoxyribonucleotides determines the order of amino acids along the polypeptide (protein) chain. The DNA sequence thus codes for the amino acid sequence.

As used herein, the term "gene" means the denxyriho- 45 nucleotide sequences comprising the coding region of a structural gene and the including sequences located adjacent to the coding region on both the 5' and 3' ends for a distance of about 1 kb on either end such that the gene corresponds to the length of the full-length mRNA. The sequences which so are located 5' of the coding region and which are present on the miRNA are referred to as 5' non-translated sequences. The sequences which are located 3' or downstream of the coding region and which are present on the mRNA are referred to as 3' non-translated sequences; these sequences. 55 The term "gene" encompasses both cDNA and genomic forms of a gene. A genomic form or clone of a gene contains the excling region interrupted with non-excling requences termed "introns" or "intervening regions" or "intervening sequences." Introns are segments of a gene which are transcribed into muclear RNA (haRNA); introns may contain regulatory elements such as enhancers. Introns are removed or "spliced out" from the nuclear or primary transcript; introns therefore are absent in the messenger RNA (mRNA) transcript. The mRNA functions during translation to 65 specify the sequence or order of amino acids in a nascent polypeptide.

In addition to containing introns, genumic forms of a gene may also include sequences located on both the 5' and 3' end of the sequences which are present on the RNA transcript. These sequences are referred to as "flanking" sequences or regions (these flanking sequences are located 5' or 3' to the non-translated sequences present on the mRNA transcript). The 5' flanking region may contain regulatory sequences such as promoters and enhancers which control or influence the transcription of the gene. The 3' flanking region may contain sequences which direct the termination of transcription, post-transcriptional cleavage and polyadeny-lation.

As used berein, the term "purified" or "to purify" refers to the removal of contaminants from a sample. For example, recombinant Cre polypeptides are expressed in bacterial host cells (as a Gst-Cre fusion protein) and the Cre polypeptides are purified by the removal of host cell proteins; the percent of recombinant Cre polypeptides is thereby increased in the sample.

The term "recombinant DNA molecule" as used herein refers to a DNA molecule which is comprised of segments of DNA joined together by means of molecular biological techniques.

The term "recombinant protein" or "recombinant polypeptide" as used herein refers to a protein molecule which is expressed from a recombinant DNA molecule.

The term "native protein" as used herein to indicate that a protein does not contain amino acid residues encoded by vector sequences; that is the native protein contains only those amino acids found in the protein as it occurs in nature. A native protein may be produced by recombinant means or may be isolated from a naturally occurring source.

the appropriate viral T antigen. Vectors which contain the replicons from bovine papillomavirus or Epstein-Barr virus protein (as in "a portion of a given protein") refers to replicate extrachromesomally at low copy number (-100 copies/cell).

As used herein, the terms "mucleic acid molecule sequence minus one amino acid.

As used herein, the term "fusion protein" refers to a chimeric protein containing the protein of interest (i.e., the Cre protein) joined to an exogenous protein fragment (the fusion partner which consists of non-Cre protein sequences). The fusion partner may enhance solubility of the protein of interest as expressed in a host cell, may provide an affinity tag to allow purification of the recombinant fusion protein from the host cell or culture supernatant, or both. If desired, the fusion protein may be removed from the pratein of interest (ie., the Cre protein) by a variety of enzymatic or chemical means known to the art.

DESCRIPTION OF THE INVENTION

The present invention provides reagents and methods which comprise a system for the rapid subcloning of nucleic acid sequences in vivo and in vitro without the need to use restriction enzymes. This system is referred to as the Univector Fusion System. The basis of the Univector Fusion System is a vector termed the Univector or the pUNI vector into which sequences encoding a gene of interest (cDNA or genomic) are inserted. The pUNI vector has a sequence-specific recombinase target site, such as a loxP site, preceding the insertion site for the gene of interest, a selectable marker gene (this feature is optional) and a conditional origin of replication that is active only in host cells expressing the requisite trans-acting replication factor. The pUNI vectors are designed to contain a gene of interest but lack a promoter for the expression of the gene of interest. The gene of interest may be cloned directly into the pUNI vector (i.e.,

the pUNI vector may be used as a cloning vector, particularly for the cloning of cDNA libraries) or a previously cloned gene of interest may be inserted (i.e., subcloned) into the pUNI vector.

Using a sequence-specific recombinase (e.g., Cre 5 recombinase), a precise fusion of the pUNI vector into a second vector containing another copy of the sequencespecific recombinase target site found on the pUNI vector is catalyzed. The second vector, referred to generically as a pHOST vector, is an expression vector that contains the 10 sequence-specific recombinase target site downstream of the promoter element contained within the expression vector. Following the site-specific recombination event which occurs between the single sequence-specific recombinase target sites located on each vector (e.g., the pUNI vector and 15 the plIOST vector), the two vectors are stably fused in a manner that places the expression of the gene of interest under the control of the promoter element contained within the expression vector. This fusion event also occurs in a manner that retains the proper translational reading frame of 20 the gene of interest. This subcloning event occurs without the need to use restriction enzymes.

The fusion or recombination event can be selected for by selecting for the ability of host cells, which do not express the trans-acting replication factor required for replication of 25 the conditional origin contained on the pUNI vector, to acquire the selectable phenotype conferred by the selectable marker gene (if present) on the pUNI vector. The pUNI vector cannot replicate in cells that do not express the trans-acting replication factor and therefore, unless the pUNI vector has integrated into the second vector that contains a non-conditional origin of replication, pUNI will be lost from the bost cell.

The Univector Pusion System allows any number of expression constructs containing the gene of interest present on the pUNI vector to be made rapidly (e.g., within a single day). Using conventional cloning or subcloning techniques which employ restriction enzyme digestion(s), the production of a single expression vector containing a gene of interest can take several days (for the design and construction of each expression vector). In contrast, once a battery of expression vectors modified to contain the appropriate sequence-specific recombinase target site is made, a gene of interest can be transferred to any number of expression vectors in an afternoon using the Univector Fusion System.

FIG. 1 provides a schematic illustrating certain elements of the pUNI vectors and the Univector Fusion System.

a) Conditional Origins of Replication and Suitable Host Cells

Conditional origins of replication are origins which require the presence or expression of a trans-acting factor in the host cell for replication. A variety of conditional origins of replication functional in prokaryotic hosts (e.g., E. coll) are known to the art. The present invention is illustrated but not limited by the use of the R6Ky origin, oriR, from the plasmid R6K. The R6Ky origin requires a trans-acting factor, the II protein supplied by the pir gene [Metcaff et al. (1996) Plasmid 35:1] E. coll strains containing the pir gene will support replication of R6Ky origins to medium copy number. A strain containing a mutant allele of pir, pir-116, will allow an even higher copy number of constructs containing the R6Ky origin.

E. coli strains that express the pir or pir-116 gene product include BW18815 (ATCC 47079; this strain contains the 65 pir-116 gene), BW19094 (AICC 47080; this strain contains the pri gene), BW20978 (this strain contains the pir-116

gene), BW20979 (this strain contains the pir gene), BW21037 (this strain contains the pir-116 gene) and BW21038 (this strain contains the pir gene) (Metcalf et al, supra).

Other conditional origins of replication suitable for use on the pUNI vectors of the present invention include:

- 1) the RK2 oriV from the plasmid RK2 (AIXX: 37125). The RK2 oriV requires a trans-acting protein encoded by the trfA gene [Ayres et al. (1993) J. Mol. Biol. 230:174];
- 2) the hacteriophage PI ori which requires the repA protein for replication [Pal et al. (1986) J. Mol. Biol. 192:275];
- 3) the origin of replication of the plasmid pSC101 (ATCC 37032) which requires a plasmid encoded protein, repA, for replication [Sugiura et al. (1992) J. Bacteriol. 175: 5993]. The pSC101 ori also requires IHF, an E. coli protein. E. coli strains carrying the himA and himD (hip) mutants (the him and hip genes encode subunits of IIII') cannot support pSC101 replication [Stenzel et al. (1987) Cell 49:709].
- 4) the bacteriophage lambda ori which requires the lambda O and P proteins [Lambda II, Hendrix et al. Eds., Cold Spring Harbor Press, Cold Spring Harbor, N.Y. (1983)];
- 5) pBR322 and other Collil derivatives will not replicate in polA mutants of *E. coli* and therefore, these origins of replication and he used in a conditional manner [Grindley and Kelley (1976) Mol. Gen. Genet. 143:311]; and
- 6) replication-thermosensitive plasmids such pSU739 or pSU300 which contain a thermosensitive replicon derived from plasmid pSC101, rep pSC101" which comprises oriV [Mendiola and de la Cruz (1989) Mol.

Microbiol. 3:979 and Francis and Lobo (1996) J. Bact. 178:894]. pSU739 and pSU300 are stably maintained in E. coli strain DHSu (Gibox BRI.) at a growth temperature of 30° C. (42° C. in non-permissive for replication of this replicon).

Other conditional origins of replication, including other temperature sensitive replicons, are known to the art and may be employed in the vectors and methods of the present invention.

b) Sequence-Specific Recombinases And Target Recognition Sites

The precise fusion between the pUNI vector and the expression vector is catalyzed by a site-specific recombinase. Site-specific recombinases are enzymes that recognize a specific DNA site or sequence (referred to herein generically as a "sequence-specific recombinate target site") and catalyzes the recombination of DNA in relation to these sites. Site-specific recombinases are employed for the recombination of DNA in both prokaryotes and cukaryotes. Examples of site-specific recombination include 1) chromosomal rearrangements which occur in Salmonella typhimurium during phase variation, inversion of the FTP sequence during the replication of the yeast 2 μ m circle and in the rearrangement of immunoglobulin and T cell receptor genes in vertebrates, 2) integration of bacteriophages into the chromosome of prokaryotic host cells to form a lysogen and 3) transposition of mobile genetic elements (e.g., transposons) in both prokaryotes and cukaryotes. The term "site specific recombinase" refers to enzymes that recognize short DNA sequences that become the crossover regions during the recombination event and includes recombinases. transposases and integrases.

The present invention is illustrated but not limited by the use of vectors containing loxP sites and the recombination of

these vectors using the Cre recombinase of bacteriophage PL The Cre protein catalyzes recombination of DNA between two loxP sites [Sternberg et al. (1981) Cold Spring Harbor Symp. Quant. Biol. 45:297]. The loxP sites may be present on the same DNA molecule or they may be present on different DNA molecules; the DNA molecules may be linear or circular or a combination of both. The loxP site consists of a double-stranded 34 bp sequence (SEQ ID NO:12) which comprises two 13 bp inverted repeat sequences separated by an 8 hp spacer region [Hoess et al. (1982) Pmc. Natl. Acad. Sci. USA 79:3398 and U.S. Pat. No. 4,959,317, the disclosure of which is herein incorporated by reference]. The internal spacer sequence of the loxP site is asymmetrical and thus, two loxP sites can exhibit directionality relative to one another [Hoese et al. (1984) Proc. Natl. Acad. Sci. USA 81:1026]. When two loxP sites on the same DNA molecule are in a directly repeated orientation, Cre excises the DNA between these two sites leaving a single lox! site on the DNA molecule [Abremski et al. (1983) Cell 32:1301]. If two loxP sites are in opposite orientation on a single DNA 20 molecule, Cre inverts the DNA sequence between these two sites rather than removing the sequence. Two circular DNA molecules each containing a single loxP site will recombine with another to form a mixture of monomer, dimer, trimer, etc. circles. The concentration of the DNA circles in the 25 reaction can be used to favor the formation of monomer (lower concentration) or multimeric circles (higher

Circular DNA molecules having a single loxP site will recombine with a linear molecule having a single loxP site 30 to produce a larger linear molecule. Cre interacts with a linear molecule containing two directly repeating loxP sites to produce a circle containing the sequences between the loxP sites and a single loxP site and a linear molecule containing a single loxP site at the site of the deletion.

The Cre protein has been purified to homogeneity [Abremski et al. (1984) J. Mol. Biol. 259:1509] and the cre gene has been cloned and expressed in a variety of host cells [Abremski et al. (1983), supra]. Purified Cre protein is available from a number of suppliers (e.g., Novagen and 40 New England Nuclear/Du Pont).

The Cre protein also recognizes a number of variant or mutant lox sites (variant relative to the loxP sequence), including the loxB, loxL and loxR sites which are found in the E. coli chromosome [Hoess et al. (1982), supra]. Other 45 Variant lox sites include loxP511 [5-ATAACTTCGTATAACTATATACGAAGTTAT-3' (SEQ ID NO:16); spacer region underlined; Hoess et al. (1986), supra], loxC2 5'-ACAAC TTCGTATA ATGTATGCTATACGAAGTTAT-3' (SEQ ID NO:17); 50 spacer region underlined; U.S. Pat. No. 4,959,317). Cre catalyzes the cleavage of the lox site within the spacer region and creates a six base-pair staggered cut [Hoesa and Abremski (1985) J. Mol. Hiol. 181:351]. The two 13 hp inverted repeat domains of the lox site represent binding sites for the 55 Cre protein. If two lox sites differ in their spacer regions in such a manner that the overhanging ends of the eleaved DNA cannot reanneal with one another, (he cannot efficiently catalyze a recombination event using the two different lox sites. For example, it has been reported that Crocannot recombine (at least not efficiently) a loxP site and a loxP511 site; these two lox sites differ in the spacer region. Two lox sites which differ due to variations in the binding sites (ie., the 13 bp inverted repeats) may be recombined by Cre provided that Cre can bind to each of the variant binding 65 sites; the efficiency of the reaction between two different lox sites (varying in the binding sites) may be less efficient that

between two lox sites having the same sequence (the efficiency will depend on the degree and the location of the variations in the binding sites). For example, the loxC2 site can be efficiently recombined with the loxP site; these two lox sites differ by a single nucleotide in the left binding site.

A variety of other site-specific recombinases may be employed in the methods of the present invention in place of the Cre recombinase. Alternative site-specific recombinases include:

- 1) the FLP recombinase of the 2pi plasmid of Saccharamyces cerevisiae [Cox (1983) Proc. Natl. Acad. Sci. USA 80:4223] which recognize the fit site which, like the loxP site, comprises two 13 bp inverted repeats separated by an 8 bp spacer [5'-GAAGTTCCTATTCTCTAGAAAGT ATAGGAACTTC-3(SEQ ID NO:18); spacer underlined]. The FLP geoe has been closed and expressed in E. coli (Cox, supra) and in mammalian cells (PCT International Patent Application PCITUS92/01899, Publication No.: WO 92/15694, the disclosure of which is herein incorporated by reference) and has been purified [Meyer-Lean et al. (1987) Nucleic Acids Res. 15:6469; Babincau et al (1985) J. Biol. Chem. 260:12313; Gronostajski and Sadowski (1985) J. Biol. Chem. 260:12328];
- the Int recombinase of bacteriophage lambda (with or without Xis) which recognizes att sites (Weisherg et al. In: Lambda II, supra, pp. 211-250);
- 3) the xerC and xerD recombinases of E. coll which together form a recombinase that recognizes the 28 bp dif site [Leslie and Sherratt (1995) EMBO J. 14:1561];
- 4) the Int protein from the conjugative transposon Tn916 [Lu and Churchward (1994) EMBO J. 13:L541];
- 5) TpnI and the β-lactamase transposons [Levesque (1990) J. Bacteriol. 172:3745];
- 6) the Tn3 resolvase [Flanagan et al. (1989) J. Mol. Biol. 206:295 and Stark et al. (1989) Cell 58:779];
- 7) the SpoIVC recombinase of Bacillus subtilis [Sato et al. J. Bacteriol. 172:1092];
- 8) the Hin recombinase [Galsgow et al. (1989) J. Biol. Chem. 264:10072];
- 9) the Cin recombinase [Hafter et al. (1988) EMHO J. 7:3991]; and
- 10) the immunoglobulin recombinases [Malynn et al. Cell (1988) 54:453].
 - c) Modification of Expression Vectors

As discussed above, pUNI vectors are used to transfer a gene of interest into a suitably modified expression vector via site-specific recombination. The modified expression vectors or host vectors used in the Univector Pusion System are referred to as pliOST vectors pliOST vectors are generally expression vectors (e.g., plasmids) which have been modified by the insertion of a sequence-specific recombinase target site (e.g., a lox site). The presence of the sequence-specific recombinase target site on the pHOST plasmid permits the rapid subcloning or insertion of the gene interest contained within a pUNI vector to generate an expression vector capable of expressing the gene of interest. The pHOST vector may encode a protein domain such as an affinity domain including, but not limited to, glutathione-Stransferase (Gst), maltose binding protein (MBP), a portion of staphylococcal protein A (SPA), a polyhistidine tract, etc. A variety of commercially available expression vectors encoding such affinity domains are known to the art. The affinity domain may be located at either the amino- or carboxy-terminus of the fusion protein. When the pHOST plasmid contains a vector-encoded affinity domain, a fusion

protein comprising the vector-encoded affinity domain and the protein of interest is generated when the pUNI and pHOST vectors are recombined.

To generate expression vectors intended to generate transcriptional fusions (i.e., pHOST does not contain a vector-sencoded protein domain), a sequence-specific recombinase target site is placed after (i.e., downstream of) the start of transcription in the bost vector; this is easily accomplished using synthetic oligonucleotides comprising the designing the sequence-specific recombinase target site. In designing the oligonucleotide comprising the sequence-specific recombinase target site, care is taken to avoid introducing an ATG or start codon that might initiate translation inappropriately.

To generate expression vectors intended to generate a fusion protein between a vector-encoded protein domain located at the amino-terminus of the fusion protein and the protein of interest (encoded by the gene of interest contained within the pUNI vector) (i.e., a translational fusion), care is taken to place the sequence-specific recombinase target site in the correct reading frame such that 1) an open reading 20 frame is maintained through the sequence-specific recombinase target site on pHOST and 2) the reading frame in the sequence-specific recombinase target site on pHOST is in frame with the reading frame found on the sequence-specific recombinase target site contained within the pUNI vector. In 25 addition, the oligonucleotide comprising the sequencespecific recombinase target site on pHOST is designed to avoid the introduction of in-frame stop codons. The gene of interest contained within the pUNI vector is cloned in a particular reading frame so as to facilitate the creation of the 30 desired fusion protein.

The modification of several expression vectors is provided in the examples below to illustrate the creation of suitable pHOST vectors. The general strategy involves the generation of a linker containing the desired sequence-specific recombinase target site (e.g., a lox site) by annealing two complementary oligonucleotides. The annealed oligonucleotides form a linker having sticky ends which are compatible with ends generated by restriction enzymes whose sites are conveniently located in the parental expression vector (e.g., within the polylinker of the parental expression vector).

d) In Vitro Recombination

The fusion of a pUNI vector and a pHOST vector may be accomplished in vitro using a purified preparation of a site-specific recombinase (e.g., Cre recombinase). The pUNI vector and the pHOST vector are placed in reaction vessel (e.g., a microcentrifuge tube) in a buffer compatible with the site-specific recombinase to be used. For example, when a Cre recombinase (native or a fusion protein form) is employed, the reaction buffer may comprise 50 mM Tris-IICI (pli 7.5), 10 mM MgCl, 30 mM NaCl and 1 mg/mi RSA. When a FLP recombinase is employed, the reaction huffer may comprise 50 mM Tris-HCl (pH 7.4), 10 mM 55 MgCl₂, 100 μg/ml BSA [Gronostajski and Sadowski (1985) 260:12320]. The concentration of the pUNI vector and the pHOST vector may vary between 100 ng to 1.0 µg of each vector per 20 pl reaction volume with about 0.1 pg of each nucleic soid construct (0.2 pg total) per 20 pl reaction being 60 preferred. The concentration of the site-specific recombinase may be titered under a standard set of reaction conditions to find the optimal concentration of enzyme to be used as described in Ex. 4.

Following the in vitro fusion reaction, a portion of the 65 reaction mixture is used to transform a suitable bost cell to permit the recovery and propagation of the fused vectors.

The best cell employed will not express the trans-acting factor required for replication of the conditional origin of replication contained within the pUNI vector (or alternatively the host cell will be grown at a temperature which is non-permissive for replication of a temperature sensitive replicon contained within the pUNI vector). The bost cells will be grown under conditions which select for the presence of the selectable marker contained within the pUNI vector (e.g., growth in the presence of kanamycin when the pUNI vector contains the kanamycin resistance gene). Plasmid or non-chromicsomal DNA is isolated from best cells which display the desired phenotype and subjected to restriction erzyme digestion to confirm that the desired fusion event has occurred.

e) Recombination in Procaryotic Host Cells

The fusion of a pUNI vector and a pHOST vector may be accomplished in vivo using a host cell that expresses the appropriate site-specific recombinase (e.g., Cre recombinase). The best cell employed will lack the shility to express the trans-acting factor required for replication of the conditional origin of replication contained within the pUNI vector (or alternatively the host cell will be grown at a temperature which is non-permissive for replication of a temperature sensitive replicon contained within the pUNI vector).

The pUNI vector and the pHOST vector are cotransformed into the host cell using a variety of methods known to the art (e.g., transformation of cells made competent by treatment with CaCl₂, electroporation, etc.). The cotransformed host cells are grown under conditions which select for the presence of the selectable marker contained within the pUNI vector (e.g., growth in the presence of kanamycin when the pUNI vector contains the kanamycin resistance gene). Plasmid or non-chromosomal DNA is isolated from host cells which display the desired phenotype and subjected to restriction enzyme digestion to confirm that the desired fusion event has occurred.

In addition to permitting the rapid transfer of a gene of interest from a particular pUNI vector containing a gene of interest into a pHOST vector, the Univector Pusion System permits the rapid exchange of an entire cDNA library to a variety of expression vectors. The cDNA library is generated using a pUNI vector as the cloning vector (a pUNI library). The entire library may then be transferred (using either an in vitro or an in vivo recombination reaction) into any expression vector modified to contain a sequence-specific recombinase target site (e.g., a lox site) (i.e., into a pHOST vector). This solves an existing problem in the art, in that there is no way, using existing vector systems, to exchange the inserts in a library made in one expression vector on masse (i.e., as an entire library) to a different expression vector. In addition, the sequences contained within a pUNI library can be used to recombine with linear λ constructs (which can then be used to isolate specific genes by complementation of an appropriate host cells such as E. coll or S. cerevisiae mutant cells). Purther as described in Example 8, the in vivo gene trap method, a variation of the Univector Pusion System, can be used to transfer linear DNA fragments that lack a selectable marker such as a PCR product into a variety of expression vectors.

EXPERIMENTAL

The following examples serve to illustrate certain preferred embodiments and aspects of the present invention and are not to be construed as limiting the scope thereof.

In the experimental disclosure which follows, the following abbreviations apply: °C. (degrees Centigrade); g

(gravitational field); vol (volume); DNA (deoxyribonucleic acid); RNA (ribonucleic acid); kdal or kD (kilodaltons); OD (optical density); EDTA (ethylene diamine tetra-acetic acid); E. coli (Escherichia coli); SDS (sodium dodecyl sulfate); PAGE (polyacrylamide gel electrophoresis); p (plasmid); 5 I.B (Luria-Bertani medium: per liter: 10 g Bacto-tryptone, 5 g yeast extract, 10 g NaCl, pll to 7.5 with NaOll); ml (milliliter); µl (microliter); M (Molar); mM (millimolar); µM (microMolar); g (gram); µg (microgram); ng (nanogram); U (units), mU (milliunits); min. (minutes); sec. (seconds); % 10 (percent); bp (base pair); kb (kilobase); PCR (polymerase chain reaction); Tris (tris(hydroxymethyl)-aminomethane); PMSF (phenylmethylsulfonylfluoride); BSA (bovine serum albumin); IPIG (isopropyl-β-D-thiogalactoside); ΛΊΧΧ (American Type Culture Collection, Rockville, Md.); His- 15 Rad (Bio-Rad Corp., Hercules, Calif.); Invitrogen (Invitrogen, Corp., San Diego, Calif.); New England Nuclear/Du Pont (Hoston, Mass.); Novagen (Novagen, Inc., Madison, Wis.); Pharmacia or Pharmacia Biotech (Pharmacia Biotech, Piscataway, N.J.); Pharmingen 20 (PharMingen, San Diegi, Calif.); Gibco BRL (Gaithersburg, Md.); and Stratagene (Stratagene Cloning Systems, La Jolla,

EXAMPLE 1

Construction Of Univector Constructs

In this example, an illustrative Univector construct is provided, the pUNI-10 vector, which contains a loxil site, a kanamyein resistance gene (Kn®) and the R6Ky conditional origin of replication (OriR_{ROEV}). The OriR_{ROEV} is functional only in E. coli strains expressing the II replication protein (i.e., the product of the pir gene). A gene of interest is placed within pUNI-10 (either as a result of constructing a library in pUNI-10 or by subcloning a previously cloned gene of 35 interest). Once the gene of interest is contained within pUNI-10, any number of plasmid expression constructs containing this gene of interest can be constructed rapidly (e.g., within a single day). The expression constructs will contain an antibiotic resistance gene other than kanamycin (e.g., ampicillin). Using the site-specific recombinase, Cre, a precise fusion between the pUNI vector and any other loxP site-containing vector comprising the desired expression signals adjacent to the loxP site is catalyzed. The sitespecific recombination event which occurs between the single loxP sites located on each plasmid (e.g., pUNI and the expression vector) results in the stable fusion of these two plasmids in such a manner as to place the expression of the gene of interest under the control of the expression signals contained within the expression vector. This subcloning event occurs without the need to use restriction enzymes. The fusion of pUNI-10 and the expression vector is selected for by selecting for the ability of E. coli cells that do not express the II protein to grow in the presence of kanamyein. pUNI cannot replicate in E. coli cells that do not express the Il protein unless pUNI has fused or integrated into another plasmid that contains a normal (i.e., not a conditional) origin of replication (e.g., the Col El origin); in this case, pUNI will he replicated (as part of the fusion plasmid) and kanamycin resistance will be conferred on the bost cell.

a) Generation of pUNI-10

FIG. 2A provides a schematic map of the pUNI-10 vector; the locations of selected restriction enzyme sites are indicated (with the exception of Notl, all sites shown are unique). FIG. 2B shows the DNA sequence of the loxP site 65 and the polylinkers contained within pUNI-10 (i.e., nucleotides 401-530 of SEQ ID NO:1).

Nucleotides 1-400 of pUNI-10 contain the conditional origin of replication from R6Ky (OriR_{MACY}); the OriR_{MACY} was derived from the plasmid R6K (ATCC 37120) [Metcalf et al. (1996) Plasmid 35:1}, nucleotides 401-414 comprise a Noti-Kpal polylinker that facilitates the exchange of lox sites; pUNI-10 contains a wild-type loxP site (as discussed above, pUNI vectors containing modified lox sites may be employed). Nucleotides 415-448 comprise the wild-type loxP site; nucleotides 449-527 comprise a polylinker used for the insertion of the gene of interest (genomic or cDNA sequences). Nucleotides 528-750 contain the polyAaddition sequence from bovine growth hormone (BGH) [the BGH polyA sequence is available on a number of commercially available vectors including pcl)NA3.1 (Invitrogen)]; the BGH polyA sequence is provides a 3' end for transcripts expressed in mammalian and other eukaryotic cells. The art is aware of other eukaryotic polyA sequences which may be used in place of the HGH polyA sequence (e.g., the SV40 poly A sequence, the TK poly A sequence, etc.). Nucleotides 751-890 contain the T7 terminator sequence which is used to terminate transcription in prokaryotic hosts (numerous prokaryotic termination signals are known to the art and may be employed in place of the 17 terminator sequence). Nucleotides 890-895 comprise an EcoRV restriction enzyme recognition site and nucleotides 896-2220 comprise the kanamycin resistance gene (Kan or Kn®) from Tn5 which provides a positive selectable marker. The Kn gene found on pUNI-10 was modified using site-directed mutagenesis to remove the naturally occurring Neol site such that pUNI-10 contains a unique Ncol site in the polylinker region located at micleotides 449-527, pUNI vectors need not contain a Kn® gene (modified or wildtype); other selectable genes may be used in place of the Kno gene (e.g., ampicillin resistance gene, tetracycline resistance gene, zeocin' resistance gene, etc.). The pUNI vector need not contain a selectable marker, although the use of a selectable marker is preferred. When a selectable marker is present on the pUNI vector, this marker is preferably a different selectable marker than that present on the pHOST vector.

The nucleotide sequence of pUNI-10 is provided in SEQ ID NO:1.

EXAMPLE 2

Construction Of Host Plasmids For Use In The Univector Plasmid-Pusion System

Host plasmids used in the Univector plasmid fusion system are referred to as pHOST plasmids, pHOST plasmids or vectors are generally expression vectors which have been modified by the insertion of a lox site. The presence of the lox site on the pHOST plasmid permits the rapid subcloning or insertion of the gene interest contained within a pUNI vector to generate an expression vector capable of expressing the gene of interest. The pHOST vector may encode a protein domain such as an affinity domain including, but not limited to, glutathione-S-transferase (Gst), maltose binding protein (MBP), a portion of staphylococcal protein A (SPA), a polyhistidine tract, etc. A variety of commercially available expression vectors encoding such affinity domains are known to the art. When the pHOST plasmid contains a vector-encoded affinity domain, a fusion protein comprising the vector-encoded affinity domain and the protein of interest is generated when the pUNI and pH()ST vectors are recombined.

To generate expression vectors intended to generate transcriptional fusions (i.e., plIOST does not contain a vector-

encoded protein domain), a lox site is placed after (i.e., downstream of) the start of transcription in the host vector; this is easily accomplished using synthetic oligonucleotides comprising the desired lox site. In designing the oligonucleotide comprising the lox site, care is taken to avoid introducing an ATG or start codon that might initiate translation inappropriately.

To generate expression vectors intended to generate a fusion protein between a vector-encoded protein domain and the protein of interest (encoded by the gene of interest contained within the pUNI vector), care is taken to place the lox site in the correct reading frame such that 1) an open reading frame is maintained through the lox site on pHOST and 2) the reading frame in the lox site on pHOST is in frame with the reading frame found on the lox site contained within the pUNI vector. In addition, the oligonucleotide comprising the lox site on pHOST is designed to avoid the introduction of in-frame stop codons. The gene of interest contained within the pUNI vector is closed in a particular reading frame so as to facilitate the creation of the desired fusion protein.

The modification of several expression vectors is provided below to illustrate the creation of suitable pHOST vectors. In each case, the general strategy involved the generation of a linker containing a lox site by annealing two complementary oligonucleotides. The armealed oligonucleotides form a linker having sticky ends which are compatible with ends generated by restriction enzymes whose sites are conveniently located in the parental expression vector (e.g., within the polylinker of the parental expression vector).

a) Modification of the pGEX-ZIKes Procaryotic Expression Vector

pGEX-ZI'Kcs is an expression vector active in 1... colicells which is designed for inducible, intracellular expression of genes or gene fragments as fusions with Gst. 35 pGEX-2TKcs contains the IPTG-inducible tac promoter (P_{tac.}) and was derived from pGEX-2TK (Pharmacia Biotech) as follows. The polylinker sequence of pGEX-2TK, 5'-GGATCCCCGGGAAITC-3' (SEQ ID NO:2), was replaced with the following sequence: 5'-GGAT CCGCATAITGCCCAAITGGCTCGAGGAITCCGAAITC-3' (SEQ ID NO:3) to generate the pGEX-2TKcs vector.

A linker containing a loxP site was generated by annealing the following oligonucleotides: 5 '-CATGGCTATAACT TCGTATAGCATACATTATGAGAGTTATG-3' (SEQ ID NO:4) and 5'-GATCCATAACTTCGTATAATGTATGC TATACGAAGTTATAGC-3' (SEQ ID NO:5). When annealed, these two oligonucleotides form a double-stranded linker having a 5' end compatible with an Nool sticky end and a 3' end compatible with a BamIII sticky end (ITIG. 3A). pGEX-2TKcs was digested with Neol and BamHII (FIG. 3B) and the annealed loxP linker was inserted to form pgst-lox.

 h) Modification of the pVI.1392 Baculovirus Expression Vector

pVL1392 is an expression vector that contains the polyhedrin promoter which is active in insect cells (Pharmingen). A linker containing a loxP site was generated by annealing the following oligonucleotides: 5'-G G C G G A C G T C A T A A C T T C G T A T A AGCATACATTATACGAAGTTATG-3' (SEQ ID NO:5) and 60 5'-GATCCATAACTTC GTATAATGTATGCTATACG AAGTTATGACGTCC-3' (SEQ ID NO:7). When annealed, these two oligonucleotides form a double-stranded linker having a 5' end compatible with a Notl sticky end and a 3' end compatible with a Notl sticky end and a 3' end compatible with a Notl sticky end (ITG. 4A). 65 pVL1392 was digested with Notl and BamHI (HG. 4B) and the annealed loxP linker was inserted to form pVL1392-lox.

c) Modification of the pGAP24 Yeast Expression Vector pGAP24 is an expression vector that is based on the yeast um circle and contains the constitutive GAP (glyceraldchyde 3-phosphate dehydrogenase) promoter (Pun) which is active in yeast cells and the TRPI gene (used a selectable marker when the cells are grown in medium lacking tryptophan) [the GAP promoter is available on pAB23; Schilds (1990) Proc. Natl. Acad. Sci. USA 87:2916]. A linker containing a loxP site was generated by annealing the following oligonucleotides: 5'-TCGAGAC GTCATÃACTTCGTAŤAGČATACATTATACGAAGTTA TGC-3' (SEQ ID NO:8) and 5'-GGCCGCATAACTTCGTA TAATGTATGCTATACGAAGTTATGACGTC-3 (SEQ ID NO:9). When annealed, these two oligonucleotides form a double-stranded linker having a 5' end compatible with a Xhol sticky end and a 3' end compatible with a Noti sticky end (FIG. 5A). pGAP24 was digested with Xhol and Notl (FIG. 5B) and the annualed loxP linker was inserted to form pGAP24-lox.

d) Modification of the pGAL14 Yeast Expression Vector pGAL14 is a yeast contromeric expression vector that contains the GAL promoter (Pall), which is induced by the presence of galactose in the medium, and the TRPI gene. A linker containing a loxP site was generated by annealing together the oligonactootides listed in SEQ II) NOS:8 and 9. When annealed, these two oligonacteotides form a double-stranded linker having a S end compatible with a XhoI sticky end and a 3' end compatible with a NotI sticky end (ITG. 6A). pGAL14 was digested with XhoI and NotI (ITG. 6B) and the annealed loxP linker was inserted to form pGAL14-lox.

EXAMPLE 3

Expression And Purification Of A Gst-Cre Fusion

In order to provide a source of purified Cre recombinase for the in vitro recombination of plasmids, the cre gene was inserted into a Gat expression vector such that a fusion protein comprising Gat at the amino-terminal end and Cre recombinase at the carboxy-terminal end was produced. The Gat-Cre fusion protein was purified by chromatography using Ghatathione Sephanose 4B (Pharmacia).

The cre gene was isolated by polymerase chain reaction (PCR) amplification using the plasmid pBS39 (U.S. Pat. No. 4,959,317). U.S. Pat. Nos. 4,663,195, 4,683,202 and 4,965, 188 cover PCR methodology and are incorporated herein by reference. The primers used in the PCR were designed to introduce an Noul site at the first ATG in the cre open reading frame. The PCR product was closed into a TA cloning vector (PCRII.1; Invitrogen) and then was subdicted as an Nool-EcoRI fragment into pGEX-2TKcs (Ex. 2) to generate pOL123. The ligation products were used to transform DH5u cells and the desired recombinant was isolated and used to transform BL21(DE3) cells (Invitrogen).

The nucleotide sequence of the Gst-Cre coding region within pQL123 is listed in SEQ ID NO:10. The amino acid sequence of the fusion protein expressed by pQL123 is listed in SEQ ID NO:11.

To express the Gst-Cre fusion protein, BL21(DE3) cells containing the pQL123 plasmid were grown at 37° C. in LB containing 100 µg/ml ampicillia until the OD 600 reached 0.6. Expression of the fusion protein was then induced by the addition of 11° 1°C to a final concentration of 0.4 mM and the cells were allowed to grow overnight at 25° C. Following

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induction, the hacterial cells were pelleted by centrifugation at 5,000 kg at 4° C. and the supernatant was discarded. A cell lysate was prepared as follows. Cells harvested from 0.5 liter of culture were suspended in 35 ml of a solution containing 20 mM Tris-IICl, pII 8.0, 0.1M NaCl, 1 mM EDTA, 0.5% 5 Nonidet P-40, 5 μ g/ml of each of leupeptin, antipain, aprotinin and 1 mM PMSF at 4° C. The cells were incubated for 10 min on ice and then disrupted by sonication (3x15 sec bursts) using a sonicator (Ultrasonic Heat Systems Model 200R) at full power. The lysate was then clarified by 10 centrifugation at 12,000 rpm using a SS34 rotor (Sorvall).

The Cist-Cre fusion protein was affinity purified from the cell lysate by chromatography on Glutathione Sepharose 4B (Pharmacia) according to the manufacturer's instructions. The protein concentration of Gst-Cre was determined by 15 Bradford analysis (BioRad).

Aliquots of the cell lysate before and after chromatography on Glutathione Sephanose 4B were applied to an SDS-PAGE gel. Following electrophoresis, the gel was stained with Coomassie blue. The stained gel is shown in FIG. 7. In FIG. 7, lanes 1 and 2 contain the cell lysate before and after chromatography, respectively. The arrowhead indicates the Gst-Cre fusion protein. The migration of the molecular weight protein markers is indicated to the left of lane 1. The results shown in FIG. 7 demonstrate the purification of the Gst-Cre fusion protein. This fusion protein was shown to be functional (i.e., capable of mediating recombination between lox sites) in the in vitro recombination assay described below.

EXAMPLE 4

In Vitro Recombination Using The Univector Plasmid Pusion System

The University Plasmid Fusion System permits the in vitro recombination of two plasmids. FIG. 8 provides a schematic showing the strategy employed for in vitro recombination. pA represents a generic pUNI vector which contains a lox1 site, a kanamycin resistance gene and the conditional R6K origin that is only functional in E. coli strains expressing the IIII protein (e.g., E. coli strains BW18815, BW19094, BW20978, BW20979, BW21037, BW21038). pB represents a generic pHOST vector which contains a lox1 site, an ampicillin resistance gene and a Col El origin of replication. pAB represents the fused plasmid which results from the Cre-mediated fusion of pA and pB.

To illustrate the in vitro recombination reaction, pUNI-5 (a pUNI vector which differs from pUNI-10 only in that 50 pUNI-5 retains the Nool site in the KoΦ gene and -contains a different polylinker) was employed as pA and pQL 103, an ampicillin-resistant plasmid containing a loxP site and the ColEl origin, was employed as pH. In a total reaction volume of 20 μl, 0.2 μg of each pUNI-5 (pA) and pQL 103 (pB) were 55 mixed in a buller containing 50 mM Tris-HCI (pH 7.5), 10 mM MgC 12, 30 mM NaC1 and 1 mg/ml BSA. The amount of purified Cist-Cre (Eix. 3) was varied from 0 to 1.0 μg. The reactions were incubated at 37° C. for 20 minutes and then the reactions were placed at 70° C. for 5 min. to inactivate 60 the GSt-Cre protein. Five microliters of each reaction mixture were used directly to transform competent DII5α cells (CaCl₂ treated). The transformed cells were plated onto LB/Amp (100 μg/ml amp) and LB/Kan (40 μg/ml kan) plates and the number of ampicillin resistant (ApΦ) and 65 kanamycin-resistant (KnΦ) colonies were counted. The results are summarized in Table 1.

TABLE 1

Gat	-Cre (pg/resution)	Ap ^a Outonies	Kn [®] Colunies	% of Thui Kn ² /Ap ²
	0	2.6 × 10 ⁴	0	0
	0.471	1.9×10^4	571	3
	0.05	1.1×10^4	682	6.2
	0.1	1.5 × 104	502	3.3
	2.0	0.3×10^4	104	34
	1.0	0.3 x 104	52	1.7

The results shown in Table 1 demonstrate, that under these reaction conditions 0.05 μ g purified fist-Cre per 20 μ l reaction yields the most efficient rate of plasmid fusion. Plasmid DNA was isolated from individual kanamycin-resistant colonics (using standard mini-prep plasmid DNA isolation protocols) and subjected to restriction enzyme digestion to determine the structure of the fused plasmids. This analysis revealed that plasmid DNA isolated from the kanamycin-resistant colonics represented a dimer created by the desired fusion of pUNI-5 and pQL103 via the loxP sites. These results demonstrate that the Univector Plasmid Fusion System can be used 20 to rapidly fuse two plasmids together in vitro.

EXAMPLE 5

In Vitro Fusion Between A pUNI Vector Containing A Gene Of Interest And A Lox-Containing Expression Vector Produces A Fused Vector Capable Of Expressing The Gene Of Interest

In Example 4 it was demonstrated that the Univector Plasmid Pusion System can be used to rapidly fuse two plasmid constructs together in vitro. In this example, the ability of the Univector Plasmid Pusion System to fuse two plasmids together in a manner that places the gene of interest contained on the pUNI vector under the transcriptional control of a promoter contained on the pHOST or expression vector in such a manner that a functional protein of interest is expressed from the fused construct.

a) Insertion Of A Gene Of Interest Into The pUNI-10 Vector

The cDNA encoding the wild-type yeast Skpl protein [Hai et al. (1996) Cell 86:263] was closed into the pUNI-10 vector between the Ndel and BamHI sites to generate pUNI-Skpl; the yeast SKP] cDNA sequence is available as GenBank accession no. U61764. Skpl is an essential protein involved in the regulation of the cell cycle in yeast. Yeast cells containing a temperature sensitive mutant of Skpl cannot grow at the non-permissive temperature (37° C.).

b) In Vitro Fusion Reactions And Complementation

pUNI-Skpl was recombined with pGAP24-lox (Ex. 2) and pGAI.14-lox (Ex. 2) using the in vitro reaction described in Ex. 4; 0.2 µg of Cbtl-Cro was used per 20 µl reaction. The resulting plasmid fusions were termed pGAP24-Skpl and pGAI.14-Skpl. pGAP24-Skpl and pGAI.14-Skpl. pGAP24-Skpl and pGAI.14-Skpl. were then transformed into the temperature sensitive (is) skpl-11 mutant yeast strain Y555 (Bai et al, supra) and the transformed yeast cells were plated onto SC-tryptophan plates (to select for the expression of the selectable marker TRPI) and incubated at either a permissive (25° C.) or non-permissive temperature (37° C.); the plates which received yeast cells transformed with pGAL14-Skpl contained galactose. The ability of the transformed cells to grow at the non-

permissive temperature is dependent upon the expression of the wild-type skpl gene encoded by a properly fused pUNI-Skpl/expression vector construct. As a control, the yeast SKPl genomic clone contained in a URA CEN vector (produced by conventional cloning techniques) was used to transform the ts skpl-11 mutant yeast strain Y555 and the transformed cells were also plated at 25° C. and 37° C. In each case, an expression vector (e.g., pRS414 or pRS415; Bai et al., supra) lacking the SKPl gene but containing the same selectable marker (ie., TRPl) as either pGAP24-Skpl, pGA1.14-Skpl or URA CEN-Skpl was used to transform Y555 cells as a control capable of permitting the growth of transformed Y555 cells on selective medium at the permissive temperature.

The results demonstrated that the URA CEN-SKPI construct produced by conventional cloming techniques produced a functional Skpl protein which was capable of complementing the lethality of the SKPI-11 is mutation. More importantly, the results demonstrated that the in vitro fusion reaction that created pGAP24-Skpl and pGAI.14-Skpl produced constructs capable of producing functional Skpl; that is, Y555 cells transformed with either pGAP24-Skpl or pGAL14-Skpl were capable of growth at 37° C., a temperature at which the ts Skpl-11 protein produced by the host strain is non-functional. Expression vectors lacking the SKPI cDNA were incapable of complementing the lethality of the skpl-11 is mutation.

 c) Restriction Analysis, SDS-PAGE Analysis and Western Blot Analysis of In Vitro Fusion Reactions

pUNI-Skpl was recombined with pGst-lox (Ex. 2) using 30 the in vitro reaction described in Ex. 4; 0.2 µg of Gst-Cre was used per 20 μ l reaction. The resulting plasmid fusion was termed pGST-Skpl. FIG. 9A provides a schematic showing the starting constructs and the predicted fusion construct. Five microliters of the fusion reaction mixture 35 was used transform DH5α cells as described in Ex. 4. The transformed cells were plated onto LB/Amp/Kan plates and plasmid DNA was isolated from individual Ap®Kn® colonies. The plasmid DNAs were digested with PstI followed by electrophoresis on agarose gels to examine the structure of the fused plasmids. A representative ethidium bromidestained gel is shown in FIG. 9B. In FIG. 9B, lane "M" contains DNA size markers, lanes pUNI-Skpl and pgst-lox contain the starting plasmids digested with PstI and lanes 1-12 contain plasmid DNA from individual Al®Kn® colonies digested with Pstl. Lanes marked with an "" indicate that these colonies contained a trimeric fusion plasmid that resulted from the fusion of two Gst-lox plasmids and one pUNI-Skpl plasmid. The sizes of the two Pstl fragments which result from the fusion of pUNI-Skpl and pgst-lox in kb are indicated (5.8 and 2.0 kb). The results shown in FIG. 9B demonstrate that the in vitro fusion reaction resulted in the production of the desired fused construct with high efficiency (about 83% of the plasmids in the AP@Kn@ colonies comprised the fusion of one pUNI-Skpl vector with 55 one pgst-lox vector).

Three individual Ap&Kn® colonies were picked and grown in liquid cultures which were induced with 1171X1 to examine whether the fused construct (pGst-Skpt) could produce the desired Gst-Skpt fusion protein. The cultures on were grown, induced and cell extracts were prepared as described in Ex. 6. An aliquot of the cell lysates prepared from induced and uninduced cells were electrophoresed on an SDS-PAGE gel and the gel was either stained with Coomaise blue or transferred to nitrocellulose to generate a 65 Western blot. The Western blot was probed using an anti-Skpl polyclonal antibody (the antibody was raised against

the yeast Skpl using conventional methods). The resulting Coomassie-stained gel and Western hlot are shown in FIGS. 10A and 10B, respectively.

In FIG. 10A, lane "M" contains protein molecular weight markers (size in kd is indicated). Lanes marked "C" contain extracts prepared from F. coli containing a GST-SKPI construct made by conventional cloning [i.e., the SKPI cDNA was excised using restriction enzymes and inserted into pGEX-2TKcs (Ex. 2)]. Lanes 1-3 contain extracts from AP®Kn® cells transformed with in vitro fusion reaction mixtures. Extracts prepared from uninduced cells and IPTG induced cells are indicated by "-" and "+", respectively. The arrowheads indicate the location of the Gst-Skpl fusion proteins; the Gst-Skpl fusion product generated from the pGST-SKPI fusion construct contains 15 additional amino acids which are located between the Gst domain and the Skpl protein sequences relative to the Gst-Skpl fusion protein expressed from the conventionally constructed GST-SKPl plasmid (the additional 15 amino acids are encoded by the linker comprising the loxP site; see FIG. 3). In FIG. 10B, the lane designations are the same as described for FIG. 10A. This Western blot confirms that the bands indicated by the arrowheads in FIG. 10A represent Gst-Skpl fusion

The results shown in FIGS. 10A and 10B demonstrate that the Univector Fusion System can be used to create an expression vector which maintains the proper translational reading frame and permits the expression of a fusion protein comprising the expression vector-encoded affinity tag and the protein of interest.

The above results demonstrate that the Univector Pusion System can be used to recombine two plasmids, one containing a gene of interest but no promoter (this vector may optionally contain expression signals such as termination signals and/or polyadenylation signals) and the other containing a promoter and optionally other expression signals (e.g., splicing signals, translation initiation codons) (and optionally sequences encoding an affinity domain) but lacking a gene of interest, in vitro in such a manner that the proper translational reading frame is maintained permitting the expression of a functional protein from the fused plasmids in the host cell.

EXVWbi T 9

Construction Of An E. coli Strain That Inducibly Expresses Cre Recombinase

An h. coli strain containing a cre gene under the control of an inducible promoter, termed the QLB4 strain, was constructed as follows. The cre gene was placed under the transcriptional control of the inducible lac promoter by inserting the cre ORF into a derivative of pNN402 [Elledge et al. (1991) Proc. Natl. Acad. Sci. USA 88:1731]; pNN402 was modified to contain a lac promoter. This construct was then crossed onto lambda phage (e.g., \(\lambda\)gtll) using conventional techniques. The recombinant lambda phage carrying the lac-cre gene was integrated into the chromosome of E. coll strain JM107 to generate the QLB4 strain.

Expression of Cre recombinase was induced by growing QLB4 cells at 37° C, until an OD_{ono} of 0.6 was reached. The culture was then split into 2 parts and IPTG was added to one part to a final concentration of 0.4 mM. As a control, the BNN132 strain (ATCC 47059; Elledge et al. (1991), supray which contains the cre gene under the transcriptional control of the endogenous cre promoter was treated as described for the QLB4 strain. Cell extracts (total protein) were prepared

from all four samples (QLB4xII/IG) and BNN132xII/IG) and examined for expression of Cre recombinase by Western blotting analysis. The Western blot was probed using a rabbit polyclonal anti-Cre antibody (Novagen) as the primary antibody and a goat anti-rabbit IgG horseradish peroxidase conjugate (Amershamn) as the secondary antibody according to the manufacturer's instructions. FIG. 11 shows a Western blot containing extracts prepared from (shown left to right) BNN123 cells grown in the absence of IPTG ("C") and QLB4 cells grown in the absence of IPTG ("C") and presence of IPTG ("QLB4+"), respectively. The location of the Cre recombinase band is indicated by the arrowbead. The additional bands seen on this Western blot are due to cross-reactivity of the crude (i.e., not affinity purified) rabbit anti-Cre antibody with bacterial proteins.

Western blot analysis demonstrated that Cre protein could 15 not be detected in BNN123 cells grown in the presence or absence of IPTG. Cre protein was detected in QLB4 cells grown in the presence of IPIG, but not in the absence of IPIG, by Western blot analysis. Therefore, the expression of Cre recombinase in QLB4 cells is greatly induced by the 20 presence of 11°I'G in the growth medium. By this analysis, the expression of Cre recombinase in QLB4 cells is dependent upon the induction of the lac-cre gene by IPTG. However, more sensitive functional assays indicate that the Cre protein is expressed constitutively at very low levels in both BNN132 cells and QLB4 cells in the absence of IPTG. In these functional assays, a pUNI vector (Kn®) and a pHOST vector (Ap®) were cotransformed into QLB4 cells and the transformed cells were grown on plates containing kanamycin to select for the presence of the pUNI-pIIOST fusion plasmid. Plasmid DNA was isolated from individual kanamycin-resistant colonies and subjected to restriction enzyme digestion to examine the structure of the plasmid DNA. This analysis revealed that multiple isoforms of the plasmid fusion product were present in the plasmid DNA 35 isolated from any single kanamycin-resistant colony. While not limiting the present invention to any particular mechanism, it is believed that low level constitutive expression of Cre recombinase leads to multiple fusion events between the pUNI and pHOST vectors resulting in the production of multimeric forms (ie., trimer, tetramer, etc.) of the fused plasmid (the desired fused plasmid is a dimer formed by fusion of pUNI and pHOS1). The multimeric plasmid fusion products would be expected to be unstable due to the fact that the Cre protein is constitutively expressed

To overcome the potential problems that low level constitutive expression of the cre gene in the host cell may cause, the expression of cre can be more tightly controlled as described below. In addition the approaches described below, the pUNI and pHOST vectors can be modified as described in Example 7 and these modified vectors can be fused using a host cell that constitutively expresses the Cre motein.

The expression of Cre recombinase can be more tightly controlled by a variety of means. For example, the expression of the cre gene can be made conditional when expressing cre under the control of the lac promoter by growing the host cells in medium containing glucose. The presence of 0.2% glucose in the growth medium virtually shuts down on transcription form the lac promoter. In addition, the lac promoter can be modified to insert additional operator (o) sites which bind the lac repressor. Other tightly controlled promoters are known to the art [e.g., the T7 promoter which requires the expression of T7 RNA polymerase; these prototes are available on the pET vectors (Novagen)] and may be employed to control the expression of the cre gene.

In addition to placing the cre ORF under the control of a tightly controlled promoter, Cre expression can be tightly controlled by placing the cre gene on a plasmid containing a temperature-sensitive (ts) replicon (e.g., rep pSC101*). When the cre gene is carried on a ts replication plasmid, Cre will be expressed during the transformation of the bost cell (because the host cell containing the ts plasmid containing the cre gene was maintained at the permissive temperature) but will be absent following recombination of the pUNI and pHOST vectors when the host cell is grown at a temperature non-permissive for replication of the ts replicon.

EXAMPLE 7

In Vivo Recombination In Procaryotic Hosts Using The Univector Fusion System

As described in Example 6, cotransformation of E. coli cells expressing Cre protein (e.g., QLB4, BNN132) with a pUNI construct and a pIIOST construct (each construct containing a single lox site) results in the fusion of these two constructs in vivo. If the host cell used for the recombination reaction constitutively expresses the Cre protein, multimeric forms of the fused constructs are generated. In addition to the methods outlined above for tightly regulating the expression of the cre gene in the host cell, cells constitutively producing Cre protein can be employed with modified pUNI and pHOST vectors as described in this example. The pUM construct is modified such that two different lox sites flank the kanamycin resistance gene (the modified pUNI construct is termed pUNI-D). The two lox sites differ in their spacer regions by one or two nucleotides and for the sake of discussion the two different lox sites are referred to as "loxA" and "loxB" (e.g., loxP and loxP511; "loxb" is used in this discussion to distinguish it from the first lox site termed "loxA" and does not indicate the use of the loxB sequence found in the E. coli chromosome). Cre cannot efficiently catalyze a recombination event between a loxA site and a loxb due to the sequence changes located in the spacer regions between the Cre binding sites; bowever Cre can efficiently catalyze the recombination between two loxA sites or two loxH sites [Hoese et al. (1986) Nucleic Acids Res.14:2287]. The pHOST construct is modified such that one loxA site and one loxB site flank the selectable marker gene (the modified pHOST construct is termed pHOST-I). In this example, pHOST contains the sacB gene as the selectable marker (a negative selectable marker). The presence of the sacB gene on pHOST-D provides a means of counter-selection as cells expressing the sacB gene are killed when the cell is grown in medium comaining 5% sucrose [Gay et al. (1985) J. Bacteriol. 164:918 and (1983) J. Bacteriol. 153:1424].

FIG. 12 provides a schematic showing the strategy for in vivo recombination in a Cre-expressing host cell (e.g., QI.B4 cells) using the pUNI-D and pHOST-D constructs. Arrows are used to indicate the direction of transcription of various genes or gene segments in FIG. 12. In FIG. 12, the following abbreviations are used: Ap@ (ampicillin resistance gene); Kn@ (kanamycin resistance gene); Ori (non-conditional plasmid origin of replication); Ori@ (the R6Ky conditional origin of replication); Cre (Cre recombinase); GENIEX (gene of interest). The strategy outlined in FIG. 12 is referred to as the "in vivo gene-trap". FIG. 12 illustrates that the second lox site (loxB) in pUNI-D (relative to the design of the pUNI-10 vector) is inserted between the kanamycin resistance gene and the R6Ky conditional origin of replication.

To generate a pHOST-D construct, a commercially available expression vector containing the desired promoter (and

optionally enhancer) is modified as described in Ex. 2 to insert the loxA site downstream of the promoter (it is not necessary that a commercially available expression vector be employed as the art is well aware of methods for the generation of expression vectors). Sequences encoding the sacB gene [Gay et al. (1983) J. Bacteriol. 153:1424; Gen-Bank accession nos. X02730 and K01987] and the second lox site (loxB) are inserted downstream of the first lox site (loxA).

The pUNI-D and pHOST-D constructs are cotransformed 10 into QLB4 cells (Ex. 6) and the transformed cells are plated onto LB/Ap/Kn plates containing 5% sucrose to select for the desired recombinant. FIG. 12 illustrates the recombination events which will occur in the presence of Cre in the QLB4 cells. First pUNI-D and pHOST-D will fuse to form 15 two dimers in which two possible double cross-over events can occur. These two double cross-over events are diagrammed in ITG. 12. The double cross-over events will result in the exchange of the DNA segments that are flanked by loxA and loxB to produce the plasmids labelled "A" and 20 "B." All plamsids which contain the sacB gene (the pHOST-D, the fused plasmids and plasmid B) will be selected against by the presence of sucrose in the growth medium. The pUNI-D construct will not be able to replicate in QLB4 cells as these cells do not express the II protein required for 25 replication of the R6Ky origin. Therefore, the only construct will be maintained in QLB4 cells selected on LB/Kn containing sucrose is the desired plasmid A in which the gene of interest from pUNI-I) has been placed under the transcriptional control of the promoter located on pHOST-D.

pUNI-10 was modified to place a second lox site, comprising the lexP511 sequence (SEQ ID NO:16) between the kanamycin resistance gene and the R6Ky conditional origin of replication to create pUNI-10-D. A second lox site, comprising the loxP511 site, was inserted onto a loxP-35 containing expression plasmid (i.e., a pHOST vector) to create a pHOST-D vector. One-half of one microgram of each plasmid was cotransformed into competent QLB4 cells and an aliquot of the transformed cells were plated onto LB/Ap plates and onto LB/Ap/Kn plates containing 5% 40 sucrose and the number of colonies on each type of plate were counted. The percentage of AP@Kn@ colonies which grew on sucrose-containing plates relative to the number of Ap@ colonics was 1% (1×10³/1×10⁵). Restriction enzyme digestion of plasmid DNA isolated from individual 45 Ap@KW@ colonies which grew on sucrose-containing plates confirmed that the desired fusions had been generated. These results indicate that the in vivo gene trap method can he used to recombine a gene of interest carried on a pUNI-D vector into an expression vector using host cells that con- so stitutively express the Cre protein.

In addition to providing a means for recombining a gene of interest carried on a pUNI-D vector into an expression vector using host cells that constitutively express the Cre protein, the in vivo gene trap method provides a means to transfer a gene of interest contained on a tinear DNA molecule (e.g., a PCR product) that lacks a selectable marker into an expression vector(s). The desired PCR product is amplified using two primers, each of which encode a different lox site (a "loxA" and "loxB" site such as a koxP and loxP511 site). A pUNI vector is constructed that contains (5' to 3') a loxA site, a counter-selectable marker such as the sacB gene and a lox B site (i.e., the two different lox sites flank the counter-selectable marker). This pUNI vector also contains a conditional origin of replication and an antibiotic resistance gene as described above and in Ex. 1. The PCR product (loxA-amplified sequence-loxB) is recombined with

the modified pUNI vector (which comprises loxA-counter-selectable marker-loxB) to create a pUNI vector containing the PCR product which now lacks the counter-selectable marker; this recombination event is selected for by growing the host cells in medium which kills the host if the counter-selectable gene is expressed. The PCR product in the pUNI vector (containing 2 lox sites) can then be placed under the courtol of the desired promoter element by recombining the pUNIXCR product construct with the appropriate pHOSI-D vector.

EXAMPLE 8

The Use Of Modified LoxP Sites To Increase Expression Of The Protein Of Interest

The pUNI and pHOST constructs employed in the Univector Plasmid Fusion System were designed such that plasmid fusion results in the introduction of a lox site between the promoter and the gene of interest. LoxP sites consist of two 13 bp inverted repeats separated by an 8 bp spacer region [Hoess et al. (1982) Proc. Natl. Acad. Sci. USA 79:3398 and U.S. Pat. No. 4,959,317]. Transcripts of the gene of interest produced from a pUNI-pHOST fusion construct comprising a loxl' site may have two 13 nucleoticle perfect inverted repeats within the 5' untranslated region (UTR) which have the potential to form a stem-loop structure (this will occur in those cases where pHOST does not encode an affinity domain at the amnino-terminus of the fusion protein). It is currently believed that the ribosome scanning mechanism is the most commonly used mechanism for initiation of translation in eukaryotes (e.g., yeast and mammalian cells). Using this mechanism, the ribosome binds to the 5' cap structure of the mRNA transcript and scans downstream along the 5' UTR searching for the first ATG or translation start codon. Without limiting the present invention to any particular mechanism, it is possible that a stem-loop structure formed by the presence of a loxP sequence on the 5' UTR of the mRNA encoding the protein of interest would block or reduce the efficiency ribosome scanning and thus the translation initiation step could be impaired. There is evidence that stem-loop structures in the 5' UTR of particular mRNAs reduce the efficiency of translation in eukaryotes [see, e.g., Donahue et al. (1988) Mol. Cell. Biol. 8:2964 and Yoon et al. Genes and Dev. (1992) 6:2463]. It is noted that no evidence suggests that the presence of a stem-loop structure in the coding region (as opposed to the 5' UIR) of a transcript negatively affects its ability to be translated. It is likely that the energy of protein synthesis is sufficient to overcome secondary structures present in mRNAs. Indeed the data presented in Ex. 5 shows that a GST-SKPI fusion construct produced using the Univector Pusion System (i.e., the construct contains a loxP site between the sequences encoding the Gst and Skpl domains) produced the same level of fusion protein as did a conventional construct encoding a Gst-Skpl fusion protein which lacks the loxP sequence. Therefore, concerns over the presence of a stem-loop structure caused by the presence of a lox sequence in a transcript encoded by a pUNI-pIIOST fusion construct are limited to those constructs which do not generate fusion proteins.

If low levels of expression are observed when a gene of interest is expressed from a pUNI-pIIOST fusion constructs comprising lox sequences that comprise perfect 13 bp inverted repeats (e.g., loxP), pUNI and pIIOST constructs containing mutated loxP sequences are employed. The mutated loxP sequences comprise point mutations that create mismatches between the two 13 bp inverted repeat

sequences within the loxl' site which disrupt the formation of or reduce the stability of a stem loop structure. Specifically, two modified loxP sites were designed that have mismatches at different positions in the inverted repeats located within a loxP site. The 13 bp inverted repeats 5 are hinding sites for the Cre protein; thus, each loxP site has two binding sites for Cre. For the purpose of discussion, these two binding sites are referred to as L and R (left and right). The wild-type loxl' site is designed L(0)-R(0) wherein "0" indicates the absence of a mutation (ie., the 10 wild-type sequence). Two derivatives of the wild-type loxP sequence were designed and termed loxl'2 and loxl'3. The sequence of lox1'2 (SEQ ID NO:13), lox1'3 (SEQ ID NO:14), as well as the wild-type lox1's sequence (SEQ II) NO:12) are shown in HG. 13. LaxP2 is placed on the 15 pUNI-10 construct (in place of the wild-type loxP site) and loxP3 is placed on the pHOST construct.

LoxP2 has repeats designated 1 (3,6)-R(0) which indicates that the third and sixth mucleotides of the left repeat are mutated; thus, a mismatch is introduced at the third and sixth 20 positions between the L and R repeats of the loxP2 site. LoxP3 has repeats designated L(0)-R(9) which indicates that the ninth nucleotide on the right repeat sequence is mutated to introduce a mismatch at the ninth position between the L and R repeats of the loxP3 site. Fusion between the loxP2 25 site on the pUNI construct and the loxP3 site on the pHOST construct will generate a hybrid loxP23 site [L(3,6)-R (9)] located between the promoter and the gene of interest and a wild-type lox! site [1 (0)-R(0)] at the distal junction. Thus, the lox123 site (SEQ ID NO:15) in the 5 UTR will have 30 three mismatches distributed at positions 3, 6 and 9 between the 13 nucleotide inverted repeats which are expected to strongly destabilize the formation of the stem-loop structure. Other mutated loxP sequences suitable for disruption of the stem-loop structure will be apparent to those skilled in the 35 art; therefore, the present invention is not limited to the use of the loxP2 and loxP3 sequences for the purpose of dis-rupting stem-loop formation on the 5' UTR of transcripts produced from pUNI-pIIOST fusion constructs. The suitability of any pair of mutated lox sites for use in the

Univector Fusion system may be tested by placing one member of the pair on a pUNI vector and the other member on a pHOST construct. The two modified vectors are then recombined in vitro as described in Ex. 4 and the fusion reaction mixture is used to transform E. coli cells and the transformed cells are plated on selective medium (e.g., on I.B/Amp and I.B/Kan plates) in order to determine the efficiency of recombination between the two mutated lox sites (Ex. 4). The efficiency of recombination between the two mutated lox sites is compared to the efficiency of recombination between two wild-type lox sites. Any pair of two different mutant lox sites that recombines at a rate that is about 5% or greater than that observed using two loxP sites is a useful pair of mutated lox sites for use in avoiding the formation of a stem-loop structure on the 5' UTR of the mRNA transcribed from the pUNI/pHOST fusion construct.

It will be apparent to those skilled in the art that a similar strategy can be employed for the modification of frt sites when the FLP recombinase is employed for the recombination event. The fit site, like lox sites, contains two 13 bp inverted repeats separated by an 8 bp spacer region.

It is clear from the above that the present invention provides methods for the subcloning of nucleic said molecules that permit the rapid transfer of a target nucleic acid sequence (e.g., a gene of interest) from nucleic acid molecule to another in vitro or in vivo without the need to rely upon restriction enzyme digestions.

All publications and patents mentioned in the above specification are herein incorporated by reference. Various modifications and variations of the described method and system of the invention will be apparent to those skilled in the art without departing from the scope and spirit of the invention. Although the invention has been described in connection with specific preferred embodiments, it should he understood that the invention as claimed should not be unduly limited to such specific embodiments. Indeed, various modifications of the described modes for carrying out the invention which are obvious to those skilled in molecular biology or related fields are intended to be within the scope of the following claims.

2 4 0

SEQUENCE LISTING

(1) GIONGRAL INFORMATIONS (i i i) NUMBER OF SPOURNERS: 18 (2) UNFORMATION FOR SIZE ID NORTH (I) SEQUENCE CHARACTERISTICS: (A) HONOTTE 2220 bas (B) TYPIC models acid (C) STRANDEDNESS, (D) TOPOLOGY: date (1 1) MCH.HCH/I.R TYPIG other madels acid (A) DESCRIPTION: Alses = "DNA" (a 1) SEQUIDECK DESCRIPTION: SIQ ID NORI: AATTCTOTCA OCCOTTAAGT OTTCCTOTOT CACTGAAAAT TOCTTTGAGA GOCTCTAAGG OCTICADA TODARATE COLOUCTION TOTOCACAR COLLARACE TAAAAFTTT AAAAGCCTTA TATATTCTTT TTTTTCTTAT AAAACTTAAA ACCTTAGAGG CTATTTAAGT TOCTGATTTA TATTAATTIT ATTOTTCAAA CATGAGAGCT TAGTACGTGA AACATGAGAG

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CTTAGTACGT	TAGCCATGAG	AGCTTAGTAC	GTTAGCCATO	AGGGTTTAGT	TCGTTAAACA	300
TUAGAGGTTA	GTACCTTAAA	CATGAGAGCT	TAGTACGTGA	AACATUAUAC	CTTAGTACGT	3 0 0
ACTATCAACA	GGTTGAACTG	CTGATCAACA	GATCCTCTAC	acaeccacae	TACCATAACT	4 2 0
TEGTATAGCA	TACATTATAC	GAAGTTATCT	GUAATTCCCC	GGGCTCGAGA	ACATATOGCC	4 8 0
					TCAGCCTCGA	
					TCCTTGACCC	
					TOGGATTOTO	
					GGGGAGGATT	
					CAAAGCCCGA	
					CCTTGGGGCC	
					ATATCCCGGG	
					CGGCGTCCCG	
					AATCTCGTGA	
					TCAGAAGAAG	1080
					ACCGTAAAGC	1140
					GGTAGCCAAC	1200
GCTATOTCCT	GATAGCGGTC	CGCCACACCC	AGCCGGCCAC	AGTCGATGAA	TCCAGAAAAG	1260
CGGCCATTTT	CCACCATGAT	ATTCGGCAAG	CAGGCATCGC	CATGGGTCAC	GACGAGATCC	1320
TCGCCGTCGG	GCATGCGCGC	CTTGAGCCTG	GCGAACAGTT	соостоосос	GAGCCCCTGA	1380
TGCTCTTCGT	CCAGATCATC	CTGATCGACA	AGACCGGCTT	CCATCCGAGT	ACGTGCTCGC	1440
TCGATGCGAT	GTÍTEGETTO	GTGGTCGAAT	GGGCAGGTAG	CCGGATCAAG	COTATGCAGC	1500
CGCCGCATTG	CATCAGCCAT	GATGGATACT	TTCTCGGCAG	GAGCAAGGTG	AGATGACAGG	1560
AGATCCTGCC	CCGGCACTTC	GCCCAATAGC	AGCCAGTCCC	TTCCCGCTTC	AGTGACAACG	1620
TCGAGCACAG	CTGCGCAAGG	AACGCCCGTC	GTGGCCAGCC	ACGATAGCCG	COCTGCCTCO	1680
					cosacocccc	1740
					TOCCCAGTCA	1 8 0 0
					ATCTTGTTCA	1860
					GCGCCATCAG	1920
					CTTACCAGAG	1980
					GTCTAGCTAT	2040
					TTCCCTTOTC	2100
					ACTGGCTTC	2160
TACCTUTTCC	CCTTCCTTTA	OCAUCCCTTO	CUCCCTGAUT	CCTTCCCCCA	CCCTCAACCT	2220

(2) INFORMATION FOR SEQ ID NO.2:

- (i) SEQUENCE CHARACTERISTICS:
 (A) LIDETTE 16 hous pain
 (B) TYPE media and
 (C) STRANDENHESS double
 (D) TOPOLOGY House
- (1 i) MOLECULE TYPE: other stockes acid
 (A) INDICRIPTION: Above = "DNA"
- (x i) SEQUENCE DESCRIPTION: SEQ ID NO.2:

-continued

GGATCCCCGG GAATTC	16
(2) INFORMATION FOR SEQ ID NO.1:	
(i) SEQUENCE CHARACTERISTICS:	
(A) LINGTH: 36 base pairs	
(R) TYPE: eucluic acid (C) STRANDEDNESS: drabbo	•
(D) TOPOLOGY: timese	
(1 i) MOLECULE TYPE: other audicic sold (A) DESCRIPTION: /dosc = "TRNA"	
(π^{-1}) SEQUIPMOS DESCRIPTION: SEQ 1D NO.5:	
GGATCOCATA TGCCCATGGC TCGAGGATCC G	GAATTC 36
(2) INPORMATION FOR SEO ID NO4:	·
(1) SEQUENCE CHARACTERISTICS:	
(A) HONGTHE 42 burn pairs (B) TYPE: nucleic acid	
(C) STRANDEDNESS: single	
(D) TOPOLOGY: these	
(i i) MOLECULE TYPE: other stacked: sciel (A) DESCRIPTION: /dosc = "DNA"	
(a i) SEQUIDICE DESCRIPTION: SRQ ID NO:4;	
CATGGCTATA ACTTCGTATA GCATACATTA T	TACGAAGTTA TG 42
(2) INFORMATION FOR SEQ ID NO.5:	•
() SEQUENCE CHARACTERISTICS:	
(A) F.FDKTTH: 42 hum puin (R) TYPR: nucluic noid	
(C) STRANDEDNESS: deglo	
vanil (ALDEFEDMAL), (CI.)	
(i i) MOLECHTE TYPE: etter necloic solid (A) DESCRIPTION: /desc = "DNA"	
(* i) SEQUENCE DESCRIPTION: SEQ ID NO.5:	
GATCCATAAC TTCGTATAAT GTATGCTATA C	COAAGTTATA GC 42
(2) INPORMATION POR SEO ID NOsé:	
() SEQUIENCE CHARACTERISTICS:	
(A) LENGTH: 46 been pains (B) TYPE: annials acid	
(C) STRANDEDNESS; dagle	
manil MUNICHUL (CI)	
(1 i) MOLECULE TYPE: etter aucteis sold (A) DESCRIPTION: /desc = "DNA"	
(= 1) SIQUIUNCE DIGERRIPTION SHQ ID NO.66	
GUCCUGACUT CATAACTTCU TATAGCATAC A	ATTATACUAA UTTATU 46
(2) INFORMATION FOR SEQ ID NO/h	
() SEQUENCE CHARACTERISTICS:	
(A) EXPRITED 46 tupe point (B) TYPE excluir acid	
(C) STRANDIONISCE Angle	
(D) TOPON OCTA House (i i) MOLECULE TYPE other sector acid	
(A) EXOLUTION: /deez = "DNA"	
(# I) SEQUENCE DESCRIPTION: SEQ ID NO.7:	

continued

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GATCCATAAC TTCOTATAAT OTATGCTATA CGAAGTTATG ACGTCC
( 2 ) INTORMATION FOR SPQ ID NO.5:
           ( i ) SEQUENCE CHARACTERISTICS:
                      ( A ) LUNGISU 46 base pairs
( B ) TYPE musicia sold
                      ( C ) STRANDRENRSS: single
                      ( D ) TOPOLOGY: linear.
         ( i i ) MOLECULE TYPE: etter audeic seid
( A ) DESCRIPTION: /doss = "DNA"
         ( # 1 ) SEQUEDICE DESCRIPTION: SEQ ID NOISE
TCGAGACGTC ATAACTTCGT ATAGCATACA TTATACGAAG TTATGC
( 2 ) INPORMATION FOR SEQ ID NO-9:
          ( i ) SEQUENCE CHARACTERISTICS:
                     ( A ) I JONETHE 46 hase pains
( B ) TYPE: suckeis acid
                      ( C ) STRANDEDNESS: single
         ( i i ) MOLECULE TYPE: ether medicle acid
( A ) DESCRIPTION: /dosc = "DNA"
         ( \star + ) SRQUIDNCK DRSCRIPTION: SRQ ID NO:9:
GGCCGCATAA CTTCGTATAA TGTATGCTAT ACGAAGTTAT GACGTC
( 2 ) INFORMATION FOR SEQ ID NO:10:
          ( i ) SEQUENCE CHARACTERISTICS:
( A ) FONTH: 1740 been pulm
( R ) TYPE: medide soid
( C ) STRANDEDMESS: double
( D ) TOPMAKPY: Bloom
        ( 1 i ) MOI PETIT R TYPR: atter meduic acid
( A ) DESCRIPTION: /desc = "DNA"
        (ir) FEATURE
                     ( A ) NAME/KEY: CD6
( B ) LOCATION: 1,1737
         ( a i ) SECUENCE DESCRIPTION: SEQ ID NO:10:
ATG TCC CCT ATA CTA GGT TAT TGG AAA ATT AAG GGC CTT GTG CAA CCC Met Ser Pro lie Lee Gly Tyr Trp Lye (le Lye Gly Lee Val Gle Pro 1 5
        COA CTT CTT TTO GAA TAT CTT GAA GAA AAA TAT GAA GAG CAT TTO Arg Lee Lee Lee Gle Tyr Lee Gle Gle Lye Tyr Gle Gle His Lee 20
TAT UAG CUC GAT GAA UGT GAT AAA TUG CUA AAC AAA AAU TTT GAA TTO
Tyr Gle Arg Asp, Gle Gly Asp Lys Trp Arg Ass Lys Pho Gle Los
35
GUT TTG GAG TTT CCC AAT CTT CCT TAT TAT ATT GAT GGT GAT GTT AAA
Gly Los Gls Phu Pro Ass Los Pro Tyr Tyr Iis Asp Gly Asp Vol Lys
SD SS 60
TTA ACA CAU TCT ATU UCC ATC ATA CUT TAT ATA UCT GAC AAG CAC AAC Lee Thr Gin Ser Mei Ain Tie Tie Arg Tyr Tie Ain And Tyn Hie Ann GS
ATG TTG GGT GGT TGT CCA AAA GAG CGT GCA GAG ATT TCA ATG CTT GAA
Met Lee Gly Gty Cyn Pro Lyn Gla Arg Ain Gla Iie Ser Met Lee Gla
85
GGA GCG GTT TTG GAT ATT AGA TAC GGT GTT TCG AGA ATT GCA TAT AGT GLy Ale Vel Lee Aep Ile Arg Tyr Gly Vel Ser Arg Ile Ale Tyr Ser 110
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244	GAC	TTT Pbo 115	GAA	ACT	CTC L	AAA Lye	OTT V • 1 1 2 0	GAT	TTT Pbc	CTT	AGC Se:	AAG L 7 + 1 2 5	CTA	CCT Pro	GAA GI o	384
ATG Met	CTG L c • 1 3 0	A A A L , .	A T G M a t	TTC Pbc	GAA GID	GAT A s p 1 3 5	COT	TTA L	7 3 7 Cy.	CAT Hia	AAA Lys 140	ACA Tb:	TAT	TTA Lcs	AAT A.s	4 3 2
1 4 3			GTA Val		130					1 3 3					1 4 0	4 # 0
V.1	V . 1	L··	TAC Tyr	M • 1	Asp	Pro	M · t	Cy.	170	A • p	Ala	76.	Pro	L 7 6	L··	5 2 K
			AAA L y e i R O					1 # 3					190			576
Lev	Ly.	S • r	AGC S • r	Lys	Т,,	11.	A L . 200	Тгр	71.	L	GIL	205	Trp	OI a	A 1 a	624
	210		GGT GI,			2 1 5					220					672
225	3	ATE	Arg	A 1 •	3 • r 2 3 0	V • 1	GIA	3 . 7	H 1 .	M + t 2 3 5	710	Me t	A. L. +	A	L • • • 2 4 0	720
L	Thr	V . I	CAC	G 1 m 2 4 5	***	L	P z 4	A L &	250	***	٧.1	A . p	A1.	7 h r 2 5 5	S • 1	768
			CGC Arg 260					265					270			# 1 6
Ph =	5 4 7	275	CAT	The	Ттр	l. y a	2 8 0	Le s	l. e n	3	V • 1	C y s 2 8 5	ATB	Ser	Trp	8 6 4
A 1 .	2 9 G	т,р	TGC Cys	l. y a	1. e u	295	A 1 1	ATG	1. y s	Т•р	3 0 0	P T •	A1.	GIL	Fra	912
305			COC ^ * 8		310					3 1 5					3 2 0	960
			ATC	3 2 3					110					3 3 5		1008
^ * &		.,	1 3 4 0	7.0	^ - 5	<i>P • u</i>	347	3 4 5	S	***	* · ·	V . I	350	1	V• I	1056
Mei	A + E	3 5 5	ATC	A + B	l. y a	G 1 .	3 6 0	V.1	Aap	A 1 •	,G 1 y	3 4 5	A * &	A 1 •	l.ya	1104
	. , ,		41.			3 7 3					3 8 0					1152
3 8 5	GI.		AUC R.,	^.,	3 • 0	Cy.	G 1 .	A.,	11.	3 9 5	^	1	A1.	Pha	1. e u 4 0 0	1200
.,,	• • •	^''	TAT	405	T & 7	1	1. 4 6	^ · n	410	^	61.	11.	A1.	415	[•	1248
Arg	V • 1	î,.	GAT A*P 420		S = r	A	The	G A C Δ # p 4 2 5	01,	GIT	AGA	M = 1	TTA L • •	D T A	HI.	1294

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ATT	GGC	AGA	ACG	AAA	ACG	CTG	GTT	AGC	ACC	GCA	GGT	GTA	GAG	AAG	GCA	1344
	•	4 3 5		•			4 4 D					4 4 5		-,-	~ . •	
						**										1392
L c •	450	Lcv	GIJ	V. 1	Tbi	455	Les	V • 1	61.	Arg	4 6 U	116	Sar	V . i	5 + 1	
GGT	GTA	GCT	GAT	GAT	cco	AAT	AAC	TAC	CTG	TTT	TGC	caa	GTC	AGA	***	1440
465	V. 1	Ale	A · p	A + p	4 7 0	A • a	A + 2	Tyr	Lco	P h c 4 7 5	C,	Azg	V . I	Arg	L 7 a	
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A 1 2	Giy	V a L	Ala	Als	Pr.	S • r	Alt	The	S • r	GIA	L	S + r	The	Arg	Ale	1488
				4 X S					4 + 0					4 0 5		
CTG	GAA	666	ATT	TTT	OAA	GCA	ACT	CAT	CGA	TTO	ATT	TAC	OOC	CCT	AAG	1536
	01.	01,	500		0.5	A I .	101	505	A14	L		171	510	All	1.7.	
GAT	GAC	7 C T	GGT	CAG	AGA	TAC	CTG	GCC	TOG	TCT	GGA	CAC	AGT	occ	COT	1584
Aip	Asp	. S e r	GI,	01.	Arg	Tyt	1 0 T	Ala	Ттр	9 • T	017	H i e 525	5 • r	ALL	Ats	
GTC	GGA	GCC	GCG	CGA	GAT	ATG	occ	coc	GCT	GGA	OTT	TCA	ATA	cca	GAG	1632
V • 1	530	A 1 •	***	Arg	Asp	M + t	A1.	A r g	A 1 s	01,	V a 1 5 4 0	3 • 1	11.	> r •	01.	
ATC	ATG	CAA	GCT	GGT	GGC	TGG	ACC	AAT	GTA	AAT	ATT	GTC	ATG	AAC	TAT	1680
545	M + 1	410	All	017	550	ттр	Thr	A	V • 1	A + 1	11.	V • 1	Met	A . z	T y r 5 6 0	
ATC	CGT		CTG	G . T	AGT	GAA	464	000	004	A T.O.	0.7.0	cac				
				AIP		GIV			ALA							1726
				5 6 5					570					5 7 5		
	GGC		TAG													1740
~ * P	~ . ,	~ * *														

(2) INPORMATION POR SEQ ID NO:11:

- (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 579 amino miris (B) TYPE: mino solid (D) TORMACY: linear

(π) SEQUENCE DESCRIPTION: SEQ ID NO:11:

(i i) MOLECULE TYPE: particle

```
Mci Ser Fro Ile Lea Gly Tyr Trp Lyo Ile Lyo Gly Lee Vet Gla Proi
The Arg Lee Lee Con Gin Tyr Lea Gin Gio Lye Tyr Gin Gin Hin Leu 20 25 30
Tyr Gla Arg Asp Gla Gly Asp Lya Trp Arg Asa Lya Lya Pha Gla Loa
35
Oly Lee Ole Phe Pro Ass Lee Pro Tyr Tyr Lie Asp Oly Asp Val Lys
Lon The Gin See Met Ale Ite Ite Arg Tyr Ite Ala Asp Lyo Hie Asn
65
Met Law Gly Gly Cyn Prw Lyn Gla Arg Ala Gla flo Sar Mat Lan Glu
85 90
Gly Ale Vel Lee Asp tie Arg Tyr Gly Vel Ser Arg tie Ale Tyr Ser
100 105
Lys Asp the Giu Thr Len Lys Val Asp the Len Ser Lys Len Pro Giu
Met Lee Lye Met Phe Glu Asp Arg Lee Cye Hie Lye Thr Tyr Leu Asu 130
Gly Asp His Val Thr His Pro Asp Pho Met Lee Tyr Asp Ala Lee Asp
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Ϋeι	V . I	Les	т, ,	M • 1	A • p	Рт•	м. :	С,.	L + w	A • p	A (.	P 6 4	Pro	Ly.	Los
V . 1	C 7 s	Pbc	L 7 . 180	L y .	Arg	116	G I 🕶	A I a 185	I 1 c	210	G t a	[14	A . p 190	Ly.	туг
f. e. s.	1. y s	S u r 1 9 5	Sar	f. y =	Ту,	110	A 1 .	T , p	P r =	f. c a	61.	(1) 2 0 5	Tip	G (.	A 1 .
Thr	7 h e 2 1 0	GIy	GIy	GIŢ	A a p	11 1 a 2 L 5	P + 0	P r o	1. y s	N e 7	A # P 2 2 0	1	V . I	P r o	A 1 6
G i y 2 2 5	S • r	AIE	Arg	A 1 a	S + r 2 3 fi	V • 1	017	5 • r	H 1 .	M • t 2 3 5		M • 1	A1.	A • =	L • 0 2 4 0
L c .	Tbr	V. I	Hi.	G 1 a 2 4 5	A . a	Los	710	A 1 •	L 2 5 0	Pro	V . I	A . p	A (.	T b : 2 5 5	8
Asp	Gı.	V . 1	A f g 2 6 0	Lys	A	Los	Mai	A . p 2 6 5	Mot	P & 0	Arg	A . p	A F & 2 7 0	Gla	A 1 .
rh.	S • •	G 1 a 2 7 5	11: 4	Thr	тер	l. y a	M e 1 2 8 D	1. • •	1	N • •	V. I	C y a 2 8 5	^ - 5	Ser	Тгр
A .	A 1 e 2 9 0	Tip	Cys	L T. o	1	A + ± 2 + 5	A 1 2	Arg	Ly.	Trp	7 h e 3 0 0	P:•	Als	01.	7:•
G L +	A = p	Val	Arg	Aip	T y / 3 1 0	Los	Los	Tys	Los	315	A1.	A # #	GI,	Les	A 1 a 3 2 0
V-1	1. y a	The	11.	325	GIA	11 1 -	1. e e	G I y	330	1	A = =	M = 1	1. • •	335	۵، د
A * B	Ser	агу	1. e m 3 4 0	Pro	Ara	Pro	Ser	3 4 S	5.,	A	A 1 +	V . 1	350	T. e. e.	V. 1
M · t	Arg	3 5 5	114	AIE	Ly.	CIT	3 6 O	V,4 1	Asp	Ale	U I y	365	Arg	A1 •	Ly.
Gli	A 1 a	Lcu	A1 •	Pbo	GI.	3 7 5	The	Аер	Pb.	A . ,	380	Val	Arg	5 • r	L··
3 8 5	G + •	A	501	A . p	3 9 0	Cy.	91.	A # p	11.	A r g 3 9 5		1	A1.	Phu	1. a a 4 0 0
Giy	i i e	Alu	Tyr	A = = 4 0 5	T b s	Loo	Lce	A r g	1 i • 4 1 0	A 1 a	GI.	11.	A1 •	A r & 4 1 5	[•
A 7 g	V • 1	1.74	4 2 0	11.	S . T	A 7 5	The	A = p 4 2 5	GIT	GIŢ	A 7 6	M · t	1. e u 4 3 0	[] .	II I A
	01,	4 3 5	Thr	Lyı	Thr	L··	V • L 4 4 0	3 . 7	Thr	A1 .	017	V 4 1	01.	L 7 4	A 1 a
Lea	5 c r 4 5 0	Les	Gly	V + 1	Thi	L y e + 5 5	Los	V . 1	01.	Ara	1 6 0	11.	Sor	۷. ۱	5
465	V. 1	A1.	^ = p	A . p	470	^ • •	^••	т,,	1. • •	4 7 5	Сув	A + 6	V . I	A r 6	1. y a 4 8 0
^	GIY	V . I	^ I •	4 # 5	***	x . ,	Al a	The	4 9 0	G i .	1. u a	8 e r	Thr	4.5	A 1 a
L.,	GI.	G 1 7	500	73.	aı.	A1 •	The	11 1 6	Ars	L	11.	т,,	3 I 7	Ale	Ly.
A . p	A . ,	5 a s	GIP	GI.	Arg	Tyr	5 2 O	A1.	T,,	3 . :	017	2 2 2 H I *	5	A 1 .	Arg
V.1	0 1 y 5 3 0	A1.	A 1 •	A+ &	A. p	5 3 5	A1.	A 7 &	A1.	GIY	V • 1	8		* * *	G I .
545	Mei	Gin	A1.	піу	3 5 0	Тир	Thr	^	V. 1	5 5 5		V . 1	Met		T , , 5 6 0
11.	A * 6	***	1. • •	5 6 5	3 4 7	G 1 w	The	017	370	Met	V • 1	A 7 5	1. • •	1. 4 B 5 7 5	(1 to
A	41 1 -	A													

```
( 2 ) INFORMATION FOR SEQ ID NO:12:
           ( I ) SEQUENCE CHARACTERISTICS:
                       (A) LENGTH: 34 base pairs
(B) TYPE: speciale acid
(C) STRANDRINKSS: drable
                        ( D ) TOPOLOGY: Hear
         ( 1 1 ) MOLICIBLE TYPE other medale acid
( A ) DESCRIPTION: /desc = "DNA"
         ( # i ) SEQUENCE DESCRIPTION: SEQ ID NO:12:
ATAACTTCGT ATAGCATACA TTATACGAAG TTAT
                                                                                                                                                                      3 4
( 2 ) INFORMATION FOR SIQ ID NO:13:
           ( i ) SEQUENCE CHARACTERISTICS:
                       (A) LENGTH: 34 base pain
(B) TYPH: mushic acid
(C) STRANDEDRESS: double
(D) TOPOLOGY: linns
         ( 1 i ) MOHJOTHHI TYPH: atlan madeic solid
( A ) DESCRIPTION: /dosc = "DNA"
         ( x i ) SEQUENCE DESCRIPTION: SEQ ID NO:13:
ATTACCTOOT ATACCATACA TTATACGAAG TTAT
                                                                                                                                                                     3 4
( 2 ) INFORMATION FOR SEQ ID NO:14:
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                       (A) LUNGTH: 34 base pains
(B) TYPE: modulo acid
(C) STRANDFONRSS: double
(D) TOPOLOGY: tiscar
         ( i i ) MOLECULE TYPE: ettax sectoic ecid
( \( \) DESCRIPTEON: /does = "DNA"
         ( x 1 ) SIGNAPHON DISSORPTIONS SHOUD NOT 14:
ATAACTICOT ATAGCATACA TTATATGAAG TTAT
                                                                                                                                                                     3 4
( 2 ) INFORMATION FOR SEQ ID NO:15:
            ( I ) SEQUENCE CHARACTERISTICS:
                       (A) LIUNCIDE 34 has pets
(B) TYPE: sectole scial
(C) STRANDEDNESS: double
(D) TOPOLOGY: these
         ( 1 i ) MOLICTHILTYPIù etter necleic acid
( A ) DESCRIPTION:/dose = "DNA"
         ( a 1 ) SECURIOROS DICECTIFICAS: SHO ID MOETS
ATTACCTEGT ATAGCATACA TTATATGAAG TTAT
                                                                                                                                                                     3 4
( 2 ) INFORMATION FOR SEQ ID NO:16:
            ( ) SECONDICE CHARACTERISTICS:
                       ( C ) STRANDEDNESS: double
( D ) FORD DOTTE: Make the pain
( R ) TYPE: market and
( C ) STRANDEDNESS: double
( D ) FORD DOTTE Hear
         ( i i ) MOTERCITE TYPE after overlets sold
( A ) DESCRIPTION: Aless = "DNA"
         ( A I ) SEQUENCE DESCRIPTION: $80 ID NO:16
ATAACTTCGT ATATATACA TTATACGAAG TTAT
                                                                                                                                                                     3 4
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3 4

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( 2 ) INFORMATION FOR SEQ ID NO:17:
          ( 1 ) SEQUENCE CHARACTERISTICS:
                     ( A ) LENGTH: 34 base pairs
                     ( B ) TYPE swelcie seid
( C ) STRANDEDNESS: double
                     ( D ) TOPOLOGY: linux
        (11) MODIANUS TYPIS a
                     JANUE TYPE other nucleic acid
(A) DESCRIPTION: /desc = "DNA"
        ( * i ) SEQUENCE DESCRIPTION: SEQ ID NO:17:
ACAACTICGT ATAATGTATG CTATACGAAG TTAT
( 2 ) INFORMATION FOR SIGNID NOTES:
          ( i ) SEQUENCE CHARACTERISTICS:
                     (A) LENOTH: 34 been pairs
(B) TYPH: medaic acid
(C) STRANDEUNESS: doub
                     (D) TOPOLOGY: liesse
        ( i i ) MOHJCORER TYPIC offer medicic acid
( A ) DESCRIPTION: /dcse = "DNA"
        ( x 1 ) SEQUENCE DESCRIPTION: SEQ ID NO:18
GAAGTTCCTA TTCTCTAGAA AGTATAGGAA CTTC
```

We claim:

1. A pUNI vector construct comprising, in 5' to 3' operable order.

a) a conditional origin of replication;

- b) a sequence-specific recombinase target site having a 5' 35 operable order: and a 3' end; and
- c) a unique restriction enzyme site, said restriction enzyme site located adjacent to said 3' end of said sequence-specific recombinase target site.
- 2. The vector construct of claim 1 further comprising a 40 prokaryotic termination sequence located 3' to said unique restriction enzyme site.
- 3. The vector construct of claim 2, wherein said prokaryotic termination sequence is the 17 termination sequence.
- 4. The vector construct of claim 1 further comprising a 45 eukaryotic polyadenylation sequence located 3' and adjacent to said unique restriction enzyme site.
- 5. The vector construct of claim 4, wherein said polyadenylation sequence is selected from the group consisting of the bovine growth hormone polyadenylation sequence, the 50 simian virus 40 polyadenylation sequence and the Herpes simplex virus thymidine kinese polyedenylation sequence.

6. The vector construct of claim 1 further comprising a scloctable marker gene.

- 7. The vector construct of claim 6, wherein said selectable 55 marker is selected from the group consisting of kanamycin resistance gene, the ampicillin resistance gene, the tetracycline resistance gene, the chloramphenical resistance gene, the streptomycin resistance gene, the strA gene and the sacB gene.
- 8. The vector construct of claim 1, wherein said sequencespecific recombinase target site is selected from the group consisting of loxP, loxP2, loxP3, loxP23, loxP511, loxB, loxC2, loxL, loxR, loxA86, loxA117, frt, dif, and att.
- 9. The vector construct of claim 1 further comprising a 65 gene of interest inserted into said unique restriction enzyme site.

10. The vector construct of claim 1, wherein said vector construct is pUNI-10, comprising the nucleotide sequence set forth in SEQ ID NO:1.

11. A pUNI vector construct comprising, in 5' to 3'

a) a conditional origin of replication;

b) a sequence-specific recombinase target site; and

- c) a polylinker, said polylinker located adjacent to said 3' end of said sequence-specific recombinase target site. 12. A pHOST expression vector construct comprising in 5' to 3' operable order:
 - a) an origin of replication;
 - b) a promoter element; and
 - c) a sequence-specific recombinase target site.
- 13. The expression vector construct of claim 12 further comprising a selectable marker gene.
- 14. A fused expression vector construct produced by the recombination of a pUNI vector and a pHOST expression vector, said fused expression vector comprising in 5' to 3' operable order:
 - a) a promoter element;
 - b) a first sequence-specific recombinase target site having a 5' and a 3' cnd;
 - c) a gene of interest joined to said 3' end of said sequencespecific recombinase target site such that a functional translational reading frame is created;
 - d) a conditional origin of replication;
 - e) a first selectable marker gene;
 - f) a second sequence-specific recombinase target site; and

g) an origin of replication.

15. The fused expression vector construct of claim 14 further comprising a second selectable marker gene.

16. A method for the in vitro recombination of a pUNI voctor and a pHOS'I expression vector to form a fused expression vector, said method comprising:

a) providing:
i) a pUNI vector construct comprising, in 5' to 3' operable order, a conditional origin of replication, a first sequence-specific recombinase target site and a first selectable marker gene;

ii) a pHOST expression vector construct comprising, in 5' to 3' operable order, an origin of replication, a promoter element and a second sequence-specific recombinase target site; and

iii) a site-specific recombinase;

b) contacting said pUNI vector and pIIOST expression vector constructs in vitro with said site-specific recombinase under conditions such that said first and second constructs are recombined to form a single fused expression vector.

17. The method of claim 16, wherein said pUNI vector construct further comprises a gene of interest located adjacent to said 3' end of said first sequence-specific recombinase target site and before the selectable marker gene, wherein the recombination of said pUNI vector and pHOST 20 expression vector constructs places said gene of interest under the transcriptional control of said promoter element within said fused expression vector.

18. The method of claim 17, wherein said pHOST expressequence encoding an affinity domain and the recombination of said pUNI vector and pHOST exression vector constructs results in placing said gene of interest in frame with said sequence encoding said affinity domain, thereby encoding a fusion protein having the affinity domain located at either the 30 amino- or carboxy-terminus of said fusion protein.

19. The method of claim 16, wherein said pHOST vector construct further comprises a second selectable marker gene, said second selectable marker being different from said first selectable marker.

20. A method for the recombination of pUNI vector and pHOST expression vector constructs to form a single fused expression vector in a prokaryotic host, comprising:

a) providing:

- i) a pUNI vector construct comprising a conditional origin of replication, a first sequence-specific recombinase target site having a 5' and a 3' end, a unique restriction enzyme site, said restriction enzyme site located adjacent to said 3' end of said sequencespecific recombinase target site, and a first selectable marker gene;
- ii) a pHOST expression vector construct comprising in 5' to 3' operable order, an origin of replication, a promoter element having a 5' and a 3' end and a second sequence-specific recombinase target site having a 5' and a 3' end;
- iii) a prokarvotic host cell expressing a site-specific recombinase;
- b) introducing said pUNI vector and pHOST expression vector constructs into said prokaryotic host cell, wherein said host cell expresses a site-specific recombinase, under conditions such that said pUNI vector and pHOST expression vector constructs are recombined to form a single fused expression vector.
- 21. The method of claim 20 further comprising growing sion vector construct further comprises a nucleotide 25 said host cell containing said single fused expression vector fuscal expression vector.
 - 22. The method of claim 20, wherein said pUNI vector construct further comprises a prokaryotic termination sequence.
 - 23. The method of claim 20, wherein said pUNI vector construct further comprises a gene of interest inserted intosaid unique restriction enzyme site.
 - 24. The method of claim 20, wherein said pUNI vector 35 construct further comprises a eukaryotic polyadenylation sequence.